

## HOSPITAL EPIDEMIOLOGY

Volume 10, Number 6 • June 1989

### **Special Update:**

Hospital Epidemiology: New Challenges and Controversies

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  - -skin/skin structure? -bones and joints?
- Convenient **B.I.D.** dosage-250 mg, 500 mg and 750 mg tablets

\*In vitro activity does not necessarily imply a correlation with in vivo results.
†Due to susceptible strains of indicated pathogens. See indicated organisms in Brief Summary.
CIPRO" SHOULD NOT BE USED IN CHILDREN, ADOLESCENTS, OR PREGNANT WOMEN.

A historyof hypersensitivity to ciprofloxacin is a contraindication to its use. A history of hypersensitivity to other quinolones may also contraindicate the use of ciprofloxacin.



Miles Inc. Pharmaceutical Division **400** Morgan Lane **West** Haven, CT 06516

Please see adjacent page of this advertisement for Brief Summary of Prescribing Information.



# CONVENIENT B.I.D. DOSAGE Dosage guidelines Mild/Moderate Infections\*: 500 mg q12h Severe/Complicated Infections\*: 750 mg q12h

#### **CIPRO**® TABLETS (ciprofloxacin HCI/Miles)

#### BRIEF SUMMARY

#### CONSULT PACKAGE INSERT FOR FULL PRESCRIBING INFORMATION

INDICATIONS AND USAGE

Cipro®is indicated for the treatment of infections caused by susceptible strains of the designated microorganisms in

the cumulatin issue useuw Lower Respiratory **Infections** caused by Escherichia coli Klebsiella pneumoniae, Enterobacter cloacae, Proteus mirabilis, Pseudomonas aeruginosa, Haemophilus influenzae Haemophilus parainfluenzae and Streptococcus

mirabilis, Pseudómonas aeruginosa, Haemophilus influenzae naemopinius paraimiuerizae en preumoniae Skin and Skin Structure Infections caused by Escherichia coli, Klebsiella pneumoniae, Enterobacter Cloacae, Proteus mirabilis Proteus vulgaris, Providencia stuartii. Morganella morganii Citrobacter freundii, Pseudomonas aeruginosa, Staphylococcus supahviococcus epidermidis and Streptococcus pyogeneosa Staphylococcus epidermidis and Streptococcus pyogeneosa aeruginosa Urinary Tract Infections caused by Escherichia coli Klebsiella pneumonae Enterobacter cloacae, Serratia marcescens, Proteus mirabilis, Providencia retigeri. Morganella morganii, Citrobacter diversus Citrobacter freundii, Pseudomonas aeruginosa. Staphylococcus epidermidis and Streptococcus laecalis Infectious Diarrhea caused by Escherichia coli enterotoxigenic strains), Campylobacter jejum Shigella flexneri.\* and Shigella sonnei\* when antibacterial therapy is indicated "Efficacy for thisorganism in this organ system was studied in fewer than 10 infections CONTRAINDICATIONS."

CONTRAINDICATIONS

A history of hypersensitivity to ciprofloxacl' is a contraindication to its "se A history of hypersensitivity to other quinolones may also contraindicate the use of ciprofloxacin

WARNINGS

WARNINGS

CIPROFLOXACIN SHOULD NOT BE USED IN CHILOREN. AOOLESCENTS, OR PREGNANT WOMEN The oral administration of ciprofloxacin caused lameness r immature dogs Histopathological examination of the weight-bearing joints of these dogs revealed permanent lesions of the cartilage Related drugs such as nalidivic acid, cinoxacin, and norfloxacin also produced erosions of cartilage of weight-bearing joints and other signs of arthropathy immature animals of various spew (SEE ANIMAL PHARMACOLOGY SECTION IN FULL PRESCRIBING INFORMATION)

PRECAUTIONS

Concret As with other quinchage clarefloxacit may cause central perious system (CNS) stimulation, which may

General: As with other quinolones clprofloxad" may cause central nervous system (CNS) stimulation, which may lead to tremor, restlessness, lightheadedness, confusion, and rarely to hallucinations or convulsive seizures. Therefore, ciprofloxacin should be used with caution "patients with know" or suspected CNS disorders, such as sever cerebral arteriosclerosistor pelipesy, or other factors whichpredispose to seizures (SEE ADVERSE REACTIONS). Anaphylactic reactions following the first dose have bee" reported in patients receiving therapy with quinolones and reactions were accompanied by cardiovascular collapse, loss of consciousness, linging pharyngeal or facil dema dyspinea. urticaria, and itching Only a few patients had a history of hypersensitivity reaction. Anaphylactic reactions may require epinephrine and other emergency measures Ciprofloxacin should be discontinued at the first sign of hypersensitivity or allergy.

reactions may require epineprinne and other energies y inclusions objectively all energy sign of hypersensitivity or actions characterized by rash fever, eosinophilia jaundice, and hepatic necrosis with fatal outcome have beer reported rarely (less than one per million prescriptions) if patients receiving ciprofloxacin along with other drugs. The possibility that these reactions were related to ciprofloxacin cannot be excluded Ciprofloxacin should be discontinued at the first appearance of a skin rash or any sign of other hypersensitivity.

reaction
Crystals of ciprofloxacl" have bee" observed rarely in the urine of human subjects but more frequently if the urine
of laboratory animals (SEE ANIMAL PHARMACOLOGY SECTION IN FULL PRESCRIBING INFORMATION). Crystalluria
related to ciprofloxacin has been reported only rarely if mar's because human uring is usually acidic Patients
receiving ciprofloxacinshould be well hydrated, and alkalinity of the urine should be avoided. The recommended daily
dose should of the dosage regiments necessary for patients within mairment of renal function (SEE DOSAGE AND
ANDIANCE TOTAIN).

ADMINISTRATION)

ADMINISTRATION)
As with any potent drug, periodic assessment of organ system functions, including renal, hepatic, and hematopotetic function, is advisable during prolonged therapy

Drug Interactions: As with other quinolones, concurrent administration of ciprofloxacin with the ophylline may lead to
elevated plasma concentrations of theophylline and prolongation of its elimination half-life. This may result in
increased risk of theophylline-related adverse reactions if concomitant use cannot be avoided, plasma levels of
theophylline should be monitored and dosage adjustments made as appropriate.

Outprolonge including elegations are also been shown to interface with the metabolism of confision. This may

Quinolones including diprofloxacin have also been shown to interfere with the metabolism of caffeine This may lead to reduced *clearance* of caffeine and a prolongation of its plasma half-life Antacids containing magnesium bydroxide or aluminum/ydroxide may interfere with the absorption of ciproflox-acm resulting r serum and urine levels lower than dewed. concurrent administration of these agents with ciprofloxacin should be avoided

acin should be avoided
Concomitant administration of the nonsteroidal anti-inflammatory drug tenbufen with a quinolone has been reported to increase the risk of CNS stimulation and convulsive seizures.
Probeneoid interferes with the renal tubular secretion of ciprofloxacin and produces a" increase it he level of ciprofloxacin in the serum This should be considered if patients are receiving both drugs concomitantly.

As with other broad-spectrum antibiotics, prolonged use of ciprofloxacin may result in overgrowth of nonsusceptible organisms. Repeated evaluation of the patient's condition and microbial susceptibility testing is essential if superinfection occurs during therapy appropriate measures should be taken.
Information for Patients: Patients should be advised that ciprofloxacin may be take with or without meals. The preferred time of dosing is two hours after a meal. Patents should also be advised to drink fluidsliberally and "of take antacids containing magnesium or aluminum Patients should be advised that ciprofloxacin may be associated with hypersensitivity reactions, even following a single dose, and to discontinue the drug at the first sign of a skin rash or other alterior reaction.

In the allocations, even following a single dose, and to discontinue the drug at the inst sign of a similar of other allergic reaction. Ciprofloxacin may cause dizziness or lightheadedness; therefore patients should know how they react to this drug before they operate a" automobile or machinery or engage " activities requiring mental alertness or coordination. Patients should be advised that ciprofloxaci" may increase the effects of theophylline and caffeine. Carcinogenesis, Mutagenesis, Impairment of Fertility-Eight in introductagenicity tests have been conducted with ciprofloxaci" and the test results are listed below. Salmonella/Microsome Test (Negative). Fertility-Eight Market and Salmonella/Microsome Test (Negative).

Salmonella/Microsome rest (Negative)

E Coli DNA Repair Assay (Negative)

MouseLymphoma Cell Forward Mutation Assay (Positive)
Chinese Hamsler V<sub>32</sub> Cell HOPRT Test (Negative)
Syrian Hamster Embryo Cell Transformation Assay (Negative)
Saccharomyces cerevisiae Point Mutation Assay (Negative)
Saccharomyces Cerevisiae Mixto Crossover and Gene Conversion Assay (Negative)

Rat Hepatocyte DNA Repair Assay (Positive)
Thus, two of the eight tests were positive, but the results of the following three *in vivo* test systems gave negative Fisher that the term tests were positive, but the results of the following three in vivo test systems gave negative results

Rat Hepatocyte DNA Repair Assay Micronicleus Test (Mice)

Dominant Lethal Test (Mice)

Long-term carcinogenicity studies "rats and mice have been completed Affer daily oral dosing for up to 2 years

there is no evidence that ciprofloxacin had any carcinogenic or tumorigenic effects if these species

Pregnancy-Pregnancy Category C: Reproduction studies have been performed in rats and mice at doses up to 6 times the usual daily human dose and have revealed no evidence of impaired fertility or harm to the fetus due to ciprofloxacin i rabbits. as with most antimicrobial agents, ciprofloxacin (30 and 100 miglkg orally) produced gastrointestinal disturbances resulting ir maternal weight loss and a rincreased incidence of abortion No teratogenicity was observed at either dose After intravenous administration at doses up to 20 mg/kg, no maternal toxicity

genicity was observed a terring dose when migration at observe to be 20 migration at observe. In addition, in internal institution was produced, and no embryotoxicity or teratogenicity was observed. There are, however, no adequate and well-controlled studies in pregnant women SINCE CIPROFLOXACIN, LIKE OTHER DRUGS IN ITS CLASS, CAUSES ARTHROPATHY IN IMMATURE ANIMALS, ITS HOULD, NOT BE USED IN PREGNANT WOMEN (SEE WARNINGS). Nursing Mothers: It is not known whether ciprofloxacin 15 excreted r' human milk; however, it is known that ciprofloxacin its excreted r' human milk. Because of this and because of the potential for senous adverse reactions from ciprofloxacin furnising infants, a decision should be made to discontinue nursing or to discontinue the drug, takinginto account the importance of the drug to the mother.

drug to the mother

Pediatric Use: Patients under the age of 18 were not included r the clinical trials of ciprofloxacin because ciprofloxacin as well as other quinolones causes arthropathy in immature animals Ciprofloxacin should not be used in
children or adolescents (SEE WARNINGS)

ADVERSE REACTIONS

ADVERSE REACTIONS
Ciprofloxacin is generally well tolerated During clinical investigation 2.799 patients received 2.868 courses of the drug Adverse events that were considered likely to be drug related on 2.799 patients received 2.868 courses of the drug Adverse events that were considered likely to be drug related on 2.799 patients received 2.868 courses, possibly related in 9.2% and remotely related in 3.0% Ciprofloxacin was discontinued because of a" adverse event 1" 3.5% of courses, primarily involving the gastrointestinal system (1.5%), kin (0.6%), and central nervous system (0.4%). Those events typical of quinolones are fathicized. The most frequently reported events, drug related or not, were nausea (5.2%) diarrhea (2.3%). vorniting (2.0%). Additional events that occurred if fess than 1% of ciprofloxacin courses are listed below. GASTROINTESTINAL: (See above) painful oral mucosa oral candidiasis dysohapia intestinal perforation, gastrointestinal bleeding. CENTRAL NERVOUS SYSTEM (See above) dizziness, lightheadedness, insomma, nightmares, hallucinations, manic reaction irritability, termor, ataxia, convulsive sezures, lethargy drowsiness, weakness: malaise, anorexia probala, depersonalization, depression, paresthesia SKINHYPERSENSITIVITY: (See above), pruritus, urticaria, photosensitivity, flushing, fever, chilis, angioedema edema of the face, neck, lips, conjunctivae or hands. cutaneous candidiasis hyperpigmentation, erythema "odasum."

'odasum.
Altergicreactions ranging from urticaria to anaphylactic reactions have been reported (SEE PRECAUTIONST SPECIAL SENSES blurred vision disturbed vision (change in color perception overbrightness of lights), decreased visual acuty, diplopia, eye pa.\*, timulus, hearing loss, bad laste MUSCULOSKELETAL joint or back pain, joint stiffness, achiness, neck or chest pain, flare-up of gout. RENALUROGENITAL interstitial nephritis, nephritis-renal failure, polyuria, urinary retention, urethral bleeding, reservice padden.

vaginitis, acuouss CARDIOVASCULAR palpitations, atrial flutter ventricular ectopy syncope, hypertension, angina pectoris, myocardial infarction, cardiopulmonary arrest cerebral thrombosis RESPIRATORY epistaxis, laryngeal or pulmonary edema, hiccough hemoptysis, dyspnea, bronchospasm, pulmonary embosism

Most of the adverse events reported were described as only mild or moderate if seventy, abated soon after the drug was discontinued, and required no treatment. In several Instances nausea vomiting, tremor, restlessness agitation, or palpitations were judged by investigations to be related to elevated plasma levels of theophyfiline possibly as a result of a drug interaction with ciprofloxacin. Other adverse events reported in the postmarketing phase include anaphyfactoid reactions. Stevens Johnson syndrome, extiplative dermatilis, toxice pidermal necrolysis, hepatic necross, postural hypotension, possible exacerbation of myasthenia gravis, confusion, dysphasia nystagmus, pseudomembranous colitis dyspepsia flatulence, and constituation Also reported were agranulocytosis elevation of serum tighteen dessaum; prolongation of prothrombin time albuminuria; candiduria, vaginal candidiasis; and renal calculi. (SEE PRECAUTIONS)

Adverse Laboratory Changes: Changes I' laboratory parameters listed as adverse events without regard to drug

niship Hepatic—Elevations of: ALT (SGPT)(19%), AST (SGOT) (17%), alkaline phosphatase (0.8%) LOH (0.4%). Serum bilirubin (0.3%) Cholestatic jaundice has bee\* reported.

Cholestatic jaundice has bee' reported.
Hematologic—eosinophilia (0 6%), leukopenia (0 4%), decreased blood platelets (0.1%). elevated blood platelets (0.1%) anarytopenia (0.18)
Renal—Elevations of Serum creatinine (1 1%). BUN (0 9%)
CRYSTALURIA CYLUNDRURIA, AND HEMATURIA HAVE BEEN REPORTED.
Other changes occurring r' less than 0.1% of courses were Elevation of serum gammaglutamyi transferase, elevation of serum amylase reduction r' blood glucose, elevated unc acid, decrease in hemoglobin, anemia, bleeding diathesis, increase in blood monocytes, and leukocytosis

VERDOSAGE

Information on curedocace in humans is not available in the prepart of earth o wordecace, the clambel should be

OVERDOSAGE
Information on overdosage in humans is not available in the event of acute overdosage, the stomach should be emptied by inducing vorniting or by gastric lavage. The patient should be carefully observed and given supportive treatment Adequate hydration must be maintained Only a small amount of ciprofloxacin (< 10%) is removed from the body after hemodialysis or peritoneal dialysis.

DOSAGE AND ADMINISTRATION

The usual adult dosage for patients with unnary tract inflections is 250 mg every 12 hours. For patients with complicated infections caused by organisms not highly susceptible. 500 mg may be administered every 12 hours. Lower respiratory tract infections, skin and skin structure infections, and bone and joint infections may be treated with 500 mg every 12 hours. For more severe or complicated infections a dosage of 750 mg may be given every 12 hours.

ours
The recommended dosage for infectious diarrhea is 500 mg every 12 hours
In patients with renal impairment, some modification of dosage is recommended (SEE DOSAGE AND ADMINIS-

In patients with erran impairment, some modification to dosage is recommended (see Dosage and Adminis-Tration Section in full prescribing information).

How Supplied

Ciprof®(ciprofloxacin HCI/Miles) is available as tablets of 250 mg 500 mg, and 750 mg " bottles of 50, and " Unit-Dose packages of 100 (SEE FULL PRESCRIBING INFORMATION FOR COMPLETE DESCRIPTION)

'Due to susceptible strains of indicated pathogens. See indicated organisms in Prescribing Information.

For further information, contact the Miles Information Service: I-800-642-4776. In VA, call collect: 703-391-7888.



#### COMMITTED TO THERAPEUTIC EFFICIENCY

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# INFECTION CONTROL

# AND HOSPITAL EPIDEMIOLOGY

EDITORIAL	Challenges and Controversies  Richard A. Garibaldi, MD; Richard P. Wenzel, MD, MSc			239
PROGRAM SUMMARIES	AIDS Risk of HIV Infection to Health Care Workers William Schaffner, MD Serologic Testing for the Human Immunodeficiency Virus— To Screen or Not to Screen			241
				243
	Dennis G. Maki, MD  Understanding the Pathogenesis of HIV Infection  Donald A. Goldmann, MD; Thomas F. Zuck, MD			248
	Expanding Roles of Hospital Epidemiology Pharmacoepidemiology John P. Burke, MD; Hugh H. Tilson, MD, MPH; Richard Platt, MD			253
	Quality Assurance Richard P. Wenzel, MD, MSc			255
	Severity of Illness Indicators Peter A. Gross, MD		257	
	Employee Health—Chemical Exposure in the Health Care Setting William A. Rutala, PhD, MPH; Bruce H. Hamory, MD  Infection Control New Problem Organisms for Infection Control John E. McGowan, Jr., MD Opportunistic Fungal Infections—The Increasing Importance of Candida Species Michael A. Pfaller, MD The Immunocompromised Host Lowell S. Young, MD			261
				267
				270
				274
	The Future of Hospital Epidemiology Calvin M. Kunin, MD			270
SPECIAL SECTIONS	<b>Topics in Clinical Microbiology:</b> Michael S. Gelfand, MD	Candida t	ropicalis	280
DEPARTMENTS	Information for Authors	238	SHEA Newsletter	285
	Calendar of Events	284	Classified Marketplace	28

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