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

Levofloxacin Tablets

Initial U.S. Approval: 1996

WARNING:

See full prescribing information for complete boxed warning

Fluoroquinolones, including level(bazacin, are associated with an increased risk of tendinitis and tendon rupture in all ages. This risk is further increased risk national general tasking corticosteroid drugs, and in patients with kidney, heart or lung transplants [See Warnings and Precautions (6.1)].
Fluoroquinolones, including level(bazacin, may exacerbate muscle weakness in persons with myasthenia gravis. Avoid level(bazacin tabletsin patients with a known history of myasthenia gravis [See Warnings and Precautions (6.2)].

To reduce the development of drug-resistant bacteria and maintain the effectiveness of levofloxacin and other antibacte drugs, k-vofloxacin tablets should be used only to treat or prevent infections that are proven or strongly suspected to be caused by bacteria, (1)

····· RECENT MAJOR CHANGES ···

- Plague (1.14)
- 04/2012
- Plague (1.14) 04/201: sage and Administration
 Dosage in Adult Patients with Normal Renal Function (2.1) 04/2012
 Dosage in Pediatric Patients (2.2) 04/2012

Dosage in Adult Patients with Normal Renal Function (2.1) 04/2012

Dosage in Padietric Patients (2.2) 04/2012

Warnings and Precautions
Macculosic letteral Discorders in Pediatric Patients and Arthropathic Effects in Animals

(4/2012)

(4/2012)

International Pressure (pseudotumor cerebra) (5.6) 10/2011

International Pressure (1.2) 10/2011

International Pressure (1.2

DOSAGE AND ADMINISTRATION Dosage in patients with normal renal function (2.1)

Type of Infection	Dose Every 24 hours	Duration (days)
Nosocomial Pneumonia (1.1)	750 mg	7-14
Community Acquired Pneumonia (1.2)	500 mg	7-14
Community Acquired Pneumonia (1.3)	750 mg	5
Acute Bacterial Sinusitis (1.4)	750 mg	5
	500 mg	10-14
Acute Bacterial Exacerbation of Chronic Bronchitis (1.5)	500 mg	7
Complicated Skin and Skin Structure Infections (SSSI) (1.6)	750 mg	7-14
Uncomplicated SSSI (1.7)	500 mg	7-10
Chronic Bacterial Prostatitis (1.8)	500 mg	28
Complicated Urinary Tract Infection (1.9) or Acute Pyelonephritis (1.11)	750 mg	5
Complicated Urinary Tract Infection (1.10) or Acute Pyelonephritis (1.11)	250 mg	10
Uncomplicated Urinary Tract Infection (1.12)	250 mg	3
Inhalational Anthrax (Post-Exposure) (1.13) Adults and		
Pediatric Patients > 50 kg	500 mg	60
Pediatric Patients < 50 kg and ≥ 6 months of age	8 mg/kg BID (not to exceed 250 mg/dose)	60
Plague (1.14)		
Adults and Pediatric Patients > 50 kg	500 mg	10 to 14
Pediatric Patients < 50 kg and ≥ 6 months of age	8 mg/kg BID (not to exceed 250 mg/dose)	10 to 14

• Adjust dose for creatinine clearance < 50 mL/min (2.3, 8.6, 12.3)

----- DOSAGE FORMS AND STRENGTHS

Formulation (3)	Strength
Tablets	250 mg, 500 mg, and 750 mg
1.00.00.00	

CONTRAINDICATIONS Known hypersensitivity to levofloxacin or other quinolones (4, 5.3)

- **Making Ambitishment of the patients and in patients with a known higher making to the patients usually over 60 years of age, in patients taking corricosteroids, and in patients with lidiney, heart or lung transplants. Discontinue if pain or inflummation in a tendon occurs (5.1,8.5)

 **May exace that muscle weakness in persons with myasthenia gravis. Avoid use in patients with a known history of 4 Annahylaries (1.5).

- myasthenia gavis (5.2).

 Anaphylatch reactions and allergic skin reactions, serious, occasionally fatal, may occur after first dose (4,5.3)

 Hematologic (including agranulocytosis, thrombocytopenia), and renal toxicities may occur after multiple doses (5.4)

 Hepatoloxicity: Severe, and sometimes fatal, hepatoxicity has been reported. Discontinue immediately if signs and symptoms of hepatitis occur (5.7)

 Central nervous system effects, including convulsions, anxiety, confusion, depression, and insomnia may occur after the first dose. Use with caution in patients with known or suspected disorders that may predispose them to sezures or lower the sezure threshold. Increased intracranial pressure (pseudotumor cerebri) has been reported (5.6)

 Clostridium difficile-associated colisis: evaluated diarriera occurs (5.7)

 Peripheral neuropathy: discontinue if symptoms occur in order to prevent irvenesibility (5.8)

 Prolongation of the QT interval and solated cases of torsadde opolites have been reported. Avoid use in patients with known prolongation, those with hypokalemia, and with other drugs that prolong the QT interval (5.9, 8.5)

----- ADVERSE REACTIONS ------

The most common reactions (23%) were nausea, headache, diarrhea, insomnia, constipation and dizziness (62).

To report SUSPECTED ADVERSE REACTIONS, contact Wockhardt USALLC., at 1-800-346-6854 or FDA at 1-800-FDA-1088 or www.fda.gov/medwarch.

DRUG INTERACTIONS

Interacting Drug	Interaction
Multivalent cation- containing products including antacids, metal cations or didanosine	Absorption of levofloxacin is decreased when the tablet formulation is taken within 2 hours of these products. (2.4, 7.1)
Warfarin	Effect may be enhanced. Monitor prothrombin time, INR, watch for bleeding (7.2)

......USE IN SPECIFIC POPULATIONS

- USE IN SPECIFIC POPULATIONS

 Geriatrics: Severe hepatotoxicity bas been reported. The majority of reports describe patients 65 years of age or older (5.5, 8.5, 17). May have increased risk of tendinopathy (including rupture), especially with concomitant corticosteroid use (5.1, 8.5, 17). May be more susceptible to prolongation of the QT interval (5.9, 8.5, 17).

 Pediatrics: Musculosise leal disorders (arthraigh, arthris), tendinopathy, and gait abnormality) seen in more tevoluscated-reacted patients than i comparature. Shown to cause arthrapshy and otsectoodnotes in piwerill animals levoluscated reacted patients than it comparature. Shown to cause arthrapshy and otsectoodnotes in piwerill animals are proportiate only for the treatment of inhalational anthrax (post-exposure) (1.13, 2.2, 8.4, 14.9) and plugue (1.14, 2.2, 8.4, 14.10)

See 17 for PATIENT COUNSELING INFORMATION and Medication Guide.

Revised: 2/2011

FULL PRESCRIBING INFORMATION: CONTENTS*

1. INDICATIONS AND USAGE

- INDICATIONS AND USAGE

 1.1 Nosocomial Preumonia
 1.2 Community-Acquired Preumonia: 7-14 day Treatment Regimen
 1.3 Community-Acquired Preumonia: 5-day Treatment Regimen
 1.3 Community-Acquired Preumonia: 5-day Treatment Regimen
 1.4 Acute Bacterial Sinusitis: 5-day and 10-14 day Treatment Regimen
 1.5 Acute Bacterial Exacerbation of Chronic Bronchitis
 1.6 Complicated Shian and Shia Structure Infections
 1.7 Uncomplicated Shian and Shia Structure Infections
 1.8 Chronic Bacterial Prostation
 1.9 Complicated Urinary Tract Infections: 10-day Treatment Regimen
 1.11 Acute Pyelonephritis: 5 or 10-day Treatment Regimen
 1.11 Acute Pyelonephritis: 5 or 10-day Treatment Regimen
 1.13 Inhalational Anthrax (Post-Exposure)
 1.13 Inhalational Anthrax (Post-Exposure)

- 1.14 Plague
 2. DOSAGE AND ADMINISTRATION
 2.1 Dosage in Adult Patients with Normal Renal Function
 2.2 Dosage in Pediatric Patients

- 2.3 Dosage Adjustment in Adults with Renal Impairment
- 2.4 Drug Interaction With Chelation Agents: A
 2.5 Administration Instructions
 3. DOSAGE FORMS AND STRENGTHS tacids, Sucralfate, Metal Cations, Multivitamins

- 3. DOSAGE FORMS AND STRENGTHS
 4. CONTRAINDICATIONS
 5. WARNINGS AND PRECAUTIONS
 5.1 Tendinopathy and Tendon Rupture
 5.2 Exacerbation of Myasthenia Gravis
 5.3 Hypersensitivity Reactions
 5.4 Other Serious and Sometimes Fatal Reactions
 5.4 University in Proceedings of the Proceeding

- 5.4 Umer Serious and Sometimes Fatal Reactions
 5.5 Hepatotixicity
 5.6 Central Nervous System Effects
 5.7 Clostridium difficile-Associated Diarrhea
 5.8 Peripheral Neuropathy
 5.9 Prolongation of the QT Interval
 5.10 Musculoskeletal Disorders in Pediatric Patients and Arthropathic Effects in Animals
 5.11 Blood Glucose Disturbances
 5.12 Photosensitivity/Phototoxicity
 5.13 Davidocoment of Drun Pasistary Bacteria
- 5.13 Development of Drug Resistant Bacteria
- 6. ADVERSE REACTIONS

- 6.1 Serious and Otherwise Important Adverse Reactions 6.2 Clinical Trial Experience
- 6.3 Postmarketing Experience
 7. DRUG INTERACTIONS
- 7. DRUG INTERÁCTIONS
 7.1 Chelation Agents: Antacids, Sucralfate, Metal Cations, Multivitamins
 7.2 Warfarin
 7.3 Antidiabetic Agents
 7.4 Non-Steroidal Anti-Inflammatory Drugs
 7.5 Theophylline
 7.6 Cyclosporine
 7.7 Digoxin
 7.7 Digoxin
 7.8 Probenecid and Cimetidine
 7.9 Interactions with Laboratory or Diagnostic Testing
 8. USE IN SPECIFIC POPULATIONS
 8.1 Pregnancy

- 8.1 Pregnancy
 8.3 Nursing Mothers
 8.4 Pediatric Use
 8.5 Geriatric Use
 8.6 Renal Impairment
 8.7 Hepatic Impairment
 10. OVERDOSAGE
 11. DESCRIPTION
 12. CLINICAL PHARM

- 12. CLINICAL PHARMACOLOGY
- 12.1 Mechanism of Action
- 12.3 Pharmacokinetics
- 12.4 Microbiology
 13. NONCLINICAL TOXICOLOGY
- 13.1 Carcinogenesis, Mutagenesis, Impairment Of Fertility 13.2 Animal Toxicology And/Or Pharmacology 14. CLINICAL STUDIES

- 14. CLINICAL STUDIES

 14.1 Nosocomial Pheumonia

 14.2 Community-Acquired Pneumonia: 7-14 day Treatment Regimen

 14.3 Community-Acquired Pneumonia: 5-day Treatment Regimen

 14.4 Acute Bacterial Sinustitis: 5-day and 10-14 day Treatment Regimen

 14.5 Complicated Skin and Skin Structure Infections

 14.6 Chronic Bacterial Prostatitis

 14.7 Complicated Urinary Tract Infections and Acute Pyelonephritis: 5-day Treatment Regimen

 14.8 Complicated Urinary Tract Infections and Acute Pyelonephritis: 10-day Treatment Regimen

 14.9 Inhalational Anthrax (Post-Exposure)

 14.10 Plague

 15. REFERENCES

 16. HOW SUPPLIED/STORAGE AND HANDLING

 16.1 Levofloxacin Tablets

 17. PATIENT COUNSELING INFORMATION

 17.1 Artibacterial Resistance

- 17.1 Artibacterial Resistance
 17.2 Administration with Food, Fluids, and Concomitant Medications
 17.3 Serious and Potentially Serious Adverse Reactions
 17.4 Drug Interactions with Insulin, Oral Hypoglycemic Agents, and Warfarin
 17.5 Plague and Anthras Studies.

- Sections or subsections omitted from the full prescribing information are not listed

FULL PRESCRIBING INFORMATION

WARNING:

Fluoroquinolones, including levofloxacin, are associated with an increased risk of tendinitis and tendon rupture in all ages. This risk is further increased in older patients usually over 60 years of age, in patients taking corticosteroid drugs, and in patients with kidney, heart or lung transplants [See Warnings and Precautions (5.1)].

Fluoroquinolones, including levofloxacin, may exacerbate muscle weakness in persons with myasthenia gravis. Avoid levofloxacin in patients with a known history of myasthenia gravis [See Warnings and Precautions (5.2)].

1. INDICATIONS AND USAGE

To reduce the development of drug-resistant bacteria and maintain the effectiveness of levofloxacin tablets and other antibacterial drugs, levofloxacin tablets 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.

Levofloxacintablets are indicated for the treatment of adults (≥18 years of age) with mild, moderate, and severe infections caused by susceptible isolates of the designated microorganisms in the conditions listed in this section.

Culture and susceptibility testing

Appropriate culture and susceptibility tests should be performed before treatment in order to isolate and identify organisms causing the infection and to determine their susceptibility to levofloxacin [see Microbiology (12-4)]. Therapy with levofloxacin tablets may be initiated before results of these tests are known; once results become available, appropriate therapy should be selected.

As with other drugs in this class, some strains of *Pseudomonas aeruginosa* may develop resistance fairly rapidly during treatment with levofloxacin tablets. Culture and susceptibility testing performed periodically during therapy will provide information about the continued susceptibility of the pathogens to the antimicrobial agent and also the possible emergence of bacterial resistance.

1.1 Nosocomial Pneumonia

Levofloxacin tablets are indicated for the treatment of nosocomial pneumonia due to methicillin-susceptible Staphylococcus aureus, Pseudomonas aeruginosa, Serratia marcescens, Escherichia coli, Klebsiella pneumoniae, Henomphilus influenzae, or Streptococcus pneumoniae, Adjunctive therapy should be used as clinically indicated. Where Pseudomonas aeruginosa is a documented or presumptive pathogen, combination therapy with an anti-pseudomonal \(\beta\)-lactam is recommended \(\left\) see \(\left\)-(Initial).

1.2 Community-Acquired Pneumonia: 7-14 day Treatment Regimen

Levofloxacin tablets are indicated for the treatment of community-acquired pneumonia due to methicillin-susceptible Staphylococcus aureus, Streptococcus pneumoniae (including multi-drug-resistant Streptococcus pneumoniae (MDRSP)), Haemophilus influenzae, Haemophilus parainfluenzae, Klebsiella pneumoniae, Moraxella catarrhalis, Chlamydophila pneumoniae, Legionella pneumophila, or Mycoplasma pneumoniae [see Dosage and Administration (2.1) and Clinical Studies (14.2)].

MDRSP isolates are strains resistant to two or more of the following antibacterials; penicillin (MIC ≥ 2 mcg/mL), 2^{md} generation cephalosporins, e.g., cefuroxime, macrolides, tetracyclines and trimethoptrimsulfamethoxazole

1.3 Community-Acquired Pneumonia: 5-day Treatment Regimen

Levofloxacintablets are indicated for the treatment of community-acquired pneumonia due to Streptococcus pneumoniae (excluding multi-drug-resistant isolates [MDRSP]), Hemophilus influenzae, Hemophilus paraillus prainfluenzae, Mycolsus pneumoniae, or Chlamydophila pneumoniae [see Dosage and Administration [2.1] and Clinical Studies (14.3). Levo floxacintablets are indicated for the treatment of acute bacterial sinusitis due to Streptococcus pneumoniae, Haemophilus influenzae, or Moraxella catarrhalis [see Clinical Studies (14.4)].

1.5 Acute Bacterial Exacerbation of Chronic Bronchitis

Levofloxacin tablets are indicated for the treatment of acute bacterial exacerbation of chronic bronchitis due to methicillin-susceptible Staphylococcus aureus, Streptococcus pneumoniae, Haemophilus influenzae, Haemophilus parainfluenzae, or Moraxella catarrhalis.

1.6 Complicated Skin and Skin Structure Infections

Levofloxacin tablets are indicated for the treatment of complicated skin and skin structure infections due to methicillin-susceptible Staphylococcus aureus, Enterococcus faecalis, Streptococcus pyogenes, or Proteus mirabilis [see Clinical Studies (14.5)].

1.7 Uncomplicated Skin and Skin Structure Infections

Levofloxacin tablets are indicated for the treatment of uncomplicated skin and skin structure infections (mild to moderate) including abscesses, cellulitis, furuncles, impetigo, pyoderma, wound infections, due to methicillin-susceptible Staphylococcus aureus, or Streptococcus pyogenes.

1.8 Chronic Bacterial Prostatitis

 $Levoflox a cin tablets are indicated for the treatment of chronic bacterial prostatitis due to \textit{Escherichia} coli, \textit{Enterococcus faecalis, or methicillin-susceptible \textit{Staphylococcus epidermidis [see Clinical Studies]} \\$

1.9 Complicated Urinary Tract Infections: 5-day Treatment Regimen

Levofloxacin tablets are indicated for the treatment of complicated urinary tract infections due to Escherichia coli, Klebsiella pneumoniae, or Proteus mirabilis [see Clinical Studies (14.7)]

1.10 Complicated Urinary Tract Infections: 10-day Treatment Regimen

Levofloxacin tablets are indicated for the treatment of complicated urinary tract infections (mild to moderate) due to Enterococcus faecalis, Enterobacter cloacae, Escherichia coli, Klebsiella pneumoniae, Proteus mirabilis, or Pseudomonas aeruginosa [see Clinical Studies (14.8)].

1.11 Acute Pyelonephritis: 5 or 10-day Treatment Regimen

Levofloxacin tablets are indicated for the treatment of acute pyelonephritis caused by Escherichia coli, including cases with concurrent bacteremia [see Clinical Studies (14.7, 14.8)].

1.12 Uncomplicated Urinary Tract Infections

Levo flox a c intablets are indicated for the treatment of uncomplicated urinary tract infections (mild to moderate) due to Escherichia coli, Klebsiella pneumoniae, or Staphylococcus saprophyticus.

1.13 Inhalational Anthrax (Post-Exposure)

1.13 Inhalational Anthrax (Post-Exposure)
Levofloxacintablets are indicated for inhalational anthrax (post-exposure) to reduce the incidence or progression of disease following exposure to aerosolized Bacillus anthracis. The effectiveness of levofloxacin is based on plasma concentrations achieved in humans, a surrogate endpoint reasonably likely to predict clinical benefit. Levofloxacinabs not been tested in humans for the post-exposure prevention of inhalation anthrax. The safety of levofloxacin in adults for durations of therapy beyond 28 days or in pediatric patients for durations of therapy beyond 14 days has not been studied. Prolonged levofloxacinherapy should only be used when the benefit outweighs the risk [see Dosage and Administration (2.1, 2.2) and Clinical Studies (14.9)].

Levofloxacintablets are indicated for treatment of plague, including pneumonic and septicemic plague, due to Yersinia pestis (Y. pestis) and prophylaxis for plague in adults and pediatric patients, 6 months of age and older. Efficacy studies of levofloxacincould not be conducted in humans with plague for ethical and feasibility reasons. Therefore, approval of this indication was based on an efficacy study conducted in animals [see Dosage and Administration (2.1, 2.2) and Clinical Studies (14.10)].

2. DOSAGE AND ADMINISTRATION

2.1 Dosage in Adult Patients with Normal Renal Function

The usual dose of levofloxacin tablets are 250 mg, 500 mg, or 750 mg administered orally every 24 hours, as indicated by infection and described in Table 1.

These recommendations apply to patients with creatinine clearance ≥ 50 mL/min. For patients with creatinine clearance <50 mL/min, adjustments to the dosing regimen are required [see Dosage and Administration (2.3)].

 $Table \ 1: Dosage \ in \ Adult \ Patients \ with \ Normal \ Renal \ Function \ (creatinine \ clearance \ge 50 \ mL/min)$

Type of Infection*	Dosed Every 24 hours	Duration (days) [†]			
Nosocomial Preumonia	750 mg	7-14			
Community Acquired Pneumonia [‡]	500 mg	7-14			
Community Acquired Pneumonia [§]	750 mg	5			
Acute Bacterial Sinusitis	750 mg	5			
	500 mg	10-14			
Acute Bacterial Exacerbation of Chronic Bronchitis	500 mg	7			
Complicated Skin and Skin Structure Infections (SSSI)	750 mg	7-14			
Uncomplicated SSSI	500 mg	7-10			
Chronic Bacterial Prostatitis	500 mg	28			
Complicated Urinary Tract Infection (cUTI) or Acute Pyelonephritis (AP)¶	750 mg	5			
Complicated Urinary Tract Infection (cUTI) or Acute Pyelonephritis (AP)#	250 mg	10			
Uncomplicated Urinary Tract Infection	250 mg	3			
Inhalational Anthrax (Post-Exposure), adult and pediatric patients > 50 kg ^{D, B}	500 mg	60 ^g			
Pediatric patients < 50 kg and ≥ 6 months of age ^{D, B}	See Table 2 below (2.2)	60 ^g			
Plague, adult and pediatric patients > 50 kg ^à	500 mg	10 to 14			
Pediatric patients < 50 kg and ≥ 6 months of age	See Table 2 below (2.2)	10 to 14			
* Due to the designated pathogens [see Indications and Usage (1)]. † Sequential therapy (intravenous to oral) may be instituted at the discretion of the physician.					
Sequential inerapy (intravenous to orai) may be instituted at the discretion of the physician. *Due to methicillin-susceptible Staphylococcus aureus, Streptococcus pneumoniae (including multi-drug-					
+Due to metati. amin-stastepione s'atapiyacoccus auteus, sueprococcus pineunoniae (including mune-mu) resistant isolates [MDRSP]), Haemophilus influenzae, Haemophilus parainfluenzae, Haemophilus par	umonhila, or Myconlasma pneumoniae [see Indications and Usage (1.2)].				
§ Due to Streptococcus pneumoniae (excluding multi-drug-resistant isolates [MDRSP]), Haemophilus influenzae, Haemophilus parainfluenzae, Mycoplasma pneumoniae, or Ch					
This regimen is indicated for cUTI due to Escherichia coli, Klebsiella pneumoniae, Proteus mirabilis and AP due to E. coli, including cases with concurrent bacteremia.	7 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1				
# This regimen is indicated for cUTI due to Enterococcus faecalis, Enterococcus cloacae, Escherichia coli, Klebsiella pneumoniae, Proteus mirabilis, Pseudomonas aeruginosa; and fu	or AP due to E. coli.				
Drug administration should begin as soon as possible after suspected or confirmed exposure to aerosolized B. anthracis. This indication is based on a surrogate endpoint. Lev	ofloxacin plasma concentrations achieved in humans are reasonably likely to predict	clinical benefit [see Clinical Studies (14.9)].			
B The safety of levofloxacin in adults for durations of therapy beyond 28 days or in pediatric patients for durations beyond 14 days has not been studied. An increased incidence of musculoskeletal adverse events compared to controls has been observed in pediatric patients [see Warnings and Precautions (5.10 Use in Specific Populations (8.4), and Clinical Studies (14.9)]. Prolonged levofloxacin therapy should only be used when the benefit outweighs the risk.					
à Drug administration should begin as soon as possible after suspected or confirmed exposure to Yersinia pestis. Higher doses of levofloxacin typically used for treatment of pne	eumonia can be used for treatment of plague, if clinically indicated.				
2.2 Dosage in Pediatric Patients					

2.2 Dosage in Pediatric Patients

The dosage in pediatric patients ≥ 6 months of age is described below in Table 2.

Table 2: Dosage in Pediatric Patients > 6 months of age

	Tuble 2. Bosuge in Fedinate Function 2 o months of age		
Type of Infection*	Dose	Freq. Once every	Duration [†]
Inhalational Anthrax (post-exposure) ^{‡,§}			
Pediatric patients > 50 kg	500 mg	24 hr	60 days§
Pediatric patients < 50 kg and ≥ 6 months of age	8 mg/kg (not to exceed 250 mg per dose)	12 hr	60 days§
Plague ¶			
Pediatric patients > 50 kg	500 mg	24 hr	10 to 14 days
Pediatric patients < 50 kg and ≥ 6 months of age	8 mg/kg (not to exceed 250 mg per dose)	12 hr	10 to 14 days

2.3 Dosage Adjustment in Adults with Renal Impairment

Administer levofloxacin with caution in the presence of renal insufficiency. Careful clinical observation and appropriate laboratory studies should be performed prior to and during therapy since elimination of levofloxacin may be reduced.

No adjustment is necessary for patients with a creatinine clearance ≥ 50 mL/min.

In patients with impaired renal function (creatinine clearance <50 mL/min), adjustment of the dosage regimen is necessary to avoid the accumulation of levofloxacin due to decreased clearance [see Use in Specific Populations (8:6)].

Table 3 shows how to adjust dose based on creatinine clearance.

Pediatric patients < 50 kg and 2 6 months of age

8 mg/kg (not to exceed 250 mg per dose)

12 hr

10 to 14 days

* Due to Bac(lines anthrack [see Indications and Usage (1.13)] and Yersinia pestis [see Indications and Usage (1.14]].

* Sequential therapy (intravenous to oral) may be instituted at the discretion of the physician.

* Drug administration should begin as soon as possible after suspected or confirmed exposure to aerosolized B. anthracks. This indication is based on a surrogate endpoint. Levofloxacin plasma concentrations achieved in humans are reasonably likely to predict clinical benefit [see Clinical Studies (14-9)]

* The safety of levofloxacin in pediatric patients for durations of therapy beyond 14 days has not been studied. An increased incidence of musculoskeletal adverse events compared to controls has been observed in pediatric patients [see Warnings and Precautions (5.10), Use in Specific Populations (6.4.), and Clinical Studies (14.9)]. Prolonged levofloxacin therapy should only be used when the benefit cutweighs the risk.

* Drug administration should begin as soon as possible after suspected or confirmed exposure to Yersinio pestis.

Dosage in Normal Renal Function Every 24 hours	Creatinine Clearance	Creatinine Clearance	Hemodialysis or Chronic Ambulatory Peritoneal Dialysis (CAPD)
	20 to 49 mL/min	10 to 19 mL/min	
750 mg	750 mg every 48 hours	750 mg initial dose, then 500 mg every 48 hours	750 mg initial dose, then 500 mg every 48 hours
500 mg	500 mg initial dose, then 250 mg every 24 hours	500 mg initial dose, then 250 mg every 48 hours	500 mg initial dose, then 250 mg every 48 hours
250 mg	No dosage adjustment required	250 mg every 48 hours. If treating uncomplicated UTI, then no dosage adjustment is required	No information on dosing adjustment is available

2.4 Drug Interaction With Chelation Agents: Antacids, Sucralfate, Metal Cations, Multivitamins

Levofloxacin Tablets should be administered at least two hours before or two hours after antacids containing magnesium, aluminum, as well as sucralfate, metal cations such as iron, and multivitamin preparations with zinc or didanosine chewable-buffered ablets or the pediatric powder for oral solution (see Drug Interactions (7.1) and Patient Counseling Information (17.2)].

2.5 Administration Instructions

Food and Levofloxacin Tablets

Levofloxacin Tablets can be administered without regard to food.

Hydration for Patients Receiving Levofloxacin Tablets

Adequate hydration of patients receiving oral levofloxacin should be maintained to prevent the formation of highly concentrated urine. Crystalluria and cylindruria have been reported with quinolones [see Adverse Reactions (6.1) and Patient Counseling Information (17.2)].

3. DOSAGE FORMS AND STRENGTHS

TABLETS, Film-coated, capsule-shaped

- 250 mg pink tablets, debossed with "W" on one side and "544" on the other side
- 500 mg peach tablets, debossed with "W" on one side and "545" on the other side
- 750 mg white tablets, debossed with "W" on one side and "547" on the other side

Levo flox acin tablets are contraindicated in persons with known hypersensitivity to levo flox acin, or other quinolone antibacterials [see Warnings and Precautions (5.3)].

5. WARNINGS AND PRECAUTIONS

5.1 Tendinopathy and Tendon Rupture

5.1 Tendinopathy and Tendon Rupture
Fluoroquinolones, including levofloxacin, are associated with an increased risk of tendinitis and tendon rupture in all ages. This adverse reaction most frequently involves the Achilles tendon, and rupture of the Achilles tendon may require surgical repair. Tendinitis and tendon rupture in the rotator cuff (the shoulder), the hand, the biceps, the thund, and other tendon sites have also been reported. The risk of developing fluoroquinolone-associated tendinitis and tendon rupture is further increased in older patients usually over 60 years of age, in those taking corticosteroid drugs, and in patients with kidney, heart or lung transplants. Factors, in addition to age and corticosteroid use, that may independently increase the risk of tendon rupture lude stremous physical activity, renal failure, and previous tendon disorders such as rheumatoid arthritis. Tendinitis and tendon rupture heave been reported in patients taking fluoroquinolones who do not have the above risk factors. Tendon rupture can occur during or after completion of therapy; cases occurring up to several months after completion of therapy have been reported. Levofloxacin should be discontinued if the patient experiences pain swelling, inflammation or rupture of a tendon. Patients should be a discontinued for the respective to the state first sign of tendinitis or tendon rupture, and to contact their healthcare provider regarding changing to a non-quinolone antimicrobial drug. [see Adverse Reactions (6.3); Patient Courseling Information (17.3)].

5.2 Exacerbation of Myasthenia Gravis

Fluoroquinolones, including levofloxacin, have neuromuscular blocking activity and may exacerbate muscle weakness in persons with myasthenia gravis. Postmarketing serious adverse events, including deaths and requirement for ventilatory support, have been associated with fluoroquinolone use in persons with myasthenia gravis. Avoid levofloxacin in patients with a known history of myasthenia gravis [see Adverse Reactions (6.3); Patient Counseling Information (17.3)].

5.3 Hypersensitivity Reactions

5.3 Hypersensitivity Reactions Serious and Constitution a

5.4 Other Serious and Sometimes Fatal Reactions

Other serious and sometimes fatal events, some due to hypersensitivity, and some due to uncertain etiology, have been reported rarely in patients receiving therapy with fluoroquimolones, including levofloxacin. These events may be severe and generally occur following the administration of multiple doses. Clinical manifestations may include one or more of the following:

• fever, rash, or severe dermatologic reactions (e.g., toxic epidermal necrolysis, Stevens-Johnson Sundrome):

- Syndrome); vasculitis; arthralgia; myalgia; serum sickness;
- allergic pneumonitis;
- interstital nephritis; acute renal insufficiency or failure; hepatitis; jaundice; acute hepatic necrosis or failure; hematis; jaundice; acute hepatic necrosis or failure; anemia, including hemolytic and aplastic; thrombocytopenia, including thromboic thrombocytopenic purpura; leukopenia; agranulocytosis; pancytopenia; and/or other hematologic abnormalities.

The drug should be discontinued immediately at the first appearance of skin rash, jaundice, or any other sign of hypersensitivity and supportive measures instituted [see Adverse Reactions (6); Patient Counseling Information (17.3)].

5.5 Hepatotoxicity

Post-marketing reports of severe hepatotoxicity (including acute hepatitis and fatal events) have been received for patients treated with levofloxacin. No evidence of serious drug-associated hepatotoxicity was detected in clinical trials of over 7,000 patients. Severe hepatotoxicity generally occurred within 14 days of initiation of therapy and most cases occurred within 6 days. Most cases of severe hepatotoxicity were not associated with hypersensitivity [see Warnings and Precoutions [6,4]]. The majority of fatal hepatotoxicity reports occurred in patients 55 years of a go or older and most were not associated with hypersensitivity. Levofloxacin should be discontinued immediately if the patient develops signs and symptoms of hepatitis [see Adverse Reactions (6); Patient Counseling Information (17.3)].

5.6 Central Nervous System Effects

Convulsions, toxic psychoses, increased intracranial pressure (including pseudotumor cerebri) have been reported in patients receiving fluoroquinolones, including levofloxacin. Fluoroquinolones may also cause central nervous system stimulation which may lead to tremors, restlessness, anxiety, also cause central nervous system stimulation which may lead to tremors, restlessness, anxiety, ightheadeness, corfusion, hallucinations, paramoia, depression, nightmares, insomnia, and, rarely, suicidal thoughts or acts. These reactions may occur following the first dose. If these reactions occur in patients receiving levofloxacin, the drug should be discontinued and appropriate measures instituted. As with other fluoroquinolnous, levofloxacin should be used with caution in patients with a known or suspected central nervous system (CNS) disorder that may predispose them to seizures or lower the seizure threshold (e.g., severe cerebral arteriosclerosis, epilepsy) or in the presence of other risk factors that may predispose them to seizures or lower the seizure threshold (e.g., certain drug therapy, renal dysfunction). [see Adverse Reactions (6): Drug Interactions (7.4, 7.5); Patient Counseling Information (17.3)].

5.7 Clostridium difficile-Associated Diarrhea

Clostridium difficile-associated diarrhea (CDAD) has been reported with use of nearly all antibacterial agents, including levofloxacin, and may range in severity from mild diarrhea to fatal colitis. 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.

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 [see Adverse Reactions (6.2), Patient Counseling Information (17.3)].

5.8 Peripheral Neuropathy

Rare cases of sensory or sensorimotor axonal polyneuropathy affecting small and/or large axons resulting in paresthesias, hypoesthesias, dysesthesias and weakness have been reported in patients receiving fluoroquinolones, including levofloxacin. Levofloxacin should be discontinued if the patient experiences symptoms of neuropathy including pain, burning, ingiling, numbness, and/or weakness or other alterations of sensation including light touch, pain, temperature, position sense, and vibratory sensation in order to prevent the development of an irreversible condition [see Adverse Reactions (6), Patient Counseling Information (17.3)].

5.9 Prolongation of the QT Interval

5.9 Prolongation of the QT Interval

Some fluoroquinolones, including levofloxacin, have been associated with prolongation of the QT interval on the electrocardiogram and infrequent cases of arrhythmia. Rare cases of torsade de pointes have been spontaneously reported during postmarketing surveillance in patients receiving fluoroquinolones, including levofloxacin. Levofloxacinshould be avoided in patients with hown prolongation of the QT interval, patients with uncorrected hypokalemia, and patients receiving Class IA (quindine, procainamide), or Class III (amiodarone, sotalol) antiarrhythmic agents. Eleterly patients may be more susceptible to drug-associated effects on the QT interval [see Adverse Reactions (6.3), Use in Specific Populations (8.5), and Patient Counseling Information (17.3)].

5.10 Mus culos keletal Disorders in Pediatric Patients and Arthropathic Effects in Animals

Levofloxacinis indicated in pediatric patients (6 months of age and older) only for the prevention of inhalational anthrax (post-exposure) and for plague [see Indications and Usage (1.13, 1.14)]. An increased incidence of musculoskeletal disorders (arthralgia, arthritis, tendinopathy, and gait abnormality) compared to controls has been observed in pediatric patients receiving levofloxacin [see Use in Specific Populations (8.4)].

In immature rats and dogs, the oral and intravenous administration of levofloxacin resulted in increased in initiative rate and copys, the oral and inaversious daministration revolutions are resulted in increase osteochondrosis. Histopathological examination of the weight-bearing joins of immature dogs dosed with levofloxacin revealed persistent lesions of the cartilage. Other fluoroquinolones also produce similar erosions in the weight-bearing joins and other signs of arthropathy in immature animals of various species [see Animal Toxicology and/or Pharmacology (13.2)].

5.11 Blood Glucose Disturbances

As with other fluoroquinolones, disturbances of blood glucose, including symptomatic hyper- and hypoglycemia, have been reported with levofloxacin, usually in diabetic patients receiving concomitant treatment with an oral hypoglycemic agent (e.g., glyburide) or with insulin. In these patients, careful monitoring of blood glucose is recommended. If a hypoglycemic reaction occurs in a patient being treated with levofloxacin, blood be discontinued and appropriate therapy should be initiated immediately [see Adverse Reactions (6.2); Drug Interactions (7.3); Patient Counseling Information (17.4)]

5.12 Photosensitivity/Phototoxicity

Moderate to severe photosensitivity/phototoxicity reactions, the latter of which may manifest as Notice are us severe promosensimylynionoxicity reactions, the fauter of winch may intimest as exaggerated surburn reactions (e.g., burning, crythema, exudation, vesicles, blistering, edema) involving areas exposed to light (typically the face, "V" area of the neck, extensor surfaces of the forearms, dorsa of the hands), can be associated with the use of fluoroquinolones after sun or UV light exposure. Therefore, excessive exposure to these sources of light should be avoided. Drug therapy should be discontinued if photosensitivity/phototoxicity occurs [see Adverse Reactions (6.3); Patient Counselling Integration (17.3). Counseling Information (17.3)].

5.13 Development of Drug Resistant Bacteria

Prescribing levofloxacin 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 [see Patient Counseling Information (17.1)].

6. ADVERSE REACTIONS

6.1 Serious and Otherwise Important Adverse Reactions

The following serious and otherwise important adverse drug reactions are discussed in greater detail in • Tendon Effects [see Warnings and Precautions (5.1)]
• Exacerbation of Ministers (2.1)

- Tendon Effects [see Warnings and Precautions (5.1)]
 Exacerbation of Myasthenia Gravis [see Warnings and Precautions (5.2)]
 Hypersensitivity Reactions [see Warnings and Precautions (5.3)]
 Other Serious and Sometimes Fatal Reactions [see Warnings and Precautions (5.4)]
 Hepatotoxicity [see Warnings and Precautions (5.5)]
 Central Nervous System Effects [see Warnings and Precautions (5.6)]
 Clostridium difficile-Associated Diarrhea [see Warnings and Precautions (5.7)]
 Perlongation of the QT Interval [see Warnings and Precautions (5.9)]
 Musculoskeletal Disorders in Pediatric Patients [see Warnings and Precautions (5.10)]
 Blood Glucose Disturbances [see Warnings and Precautions (5.11)]
 PhotosersitivityPhototoxicity [see Warnings and Precautions (5.11)]
 Development of Drug Resistant Bacteria [see Warnings and Precautions (5.13)]

Crystalluria and cylindruria have been reported with quinolones, including levofloxacin. Therefore, adequate hydration of patients receiving levofloxacinshould be maintained to prevent the formation of a highly concentrated urine [see Dosage and Administration (2.5]].

6.2 Clinical Trial Experience

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

The data described below reflect exposure to levofloxacin in 7537 patients in 29 pooled Phase 3 clinical trials. The population studied had a mean age of 50 years (approximately 74% of the population was < 65 years of age), 50% over male, 71% were Caucasian, 19% were Black. Patients were treated with levofloxacin for a wide variety of infectious diseases [see Indications and Usage (1)]. Patients received levofloxacin forse or 750 mg once of adily, 250 mg once adily, or 500 mg once or twice daily. Treatment duration was usually 3-14 days, and the mean number of days on therapy was 10 days.

Treatment duration was usuary 57-14 days, and use mean name of days of indecipy was 10 days. The overall incidence, type and distribution of adverse reactions was similar in patients receiving levofloxacin doses of 750 mg once daily, 250 mg once daily, and 500 mg once or twice daily. Discontinuation of levofloxacin due to adverse drug reactions occurred in 4.3% of patients overall, 3.8% of patients reated with the 250 mg and 500 mg doses and 5.4% of patients treated with the 750 mg dose. The most common adverse drug reactions leading to discontinuation with the 250 and 500 mg doses were gastrointestinal (1.4%), primarily muses (0.6%), vonting (0.4%), dizziness (0.3%); and headache (0.2%). The most common adverse drug reactions leading to discontinuation with the 750 mg decreases a continuation of 1000 mg doses. dose were gastrointestinal (1.2%), primarily nausea (0.6%), vomiting (0.5%); dizziness (0.3%); and headache (0.3%).

Adverse reactions occurring in ≥1% of levofloxacin-treated patients and less common adverse reactions, occurring in 0.1 to <1% of levofloxacin-treated patients, are shown in Table 4 and Table 5, respectively. The most common adverse drug reactions (≥3%) are nausea, headache, diarrhea, insomnia, constipation, and dizziness.

Table 4: Common (≥1%) Adverse Reactions Reported in Clinical Trials with Levofloxacin

System/Organ Class	Adverse Reaction	% (N=7537)
Infections and Infestations	moniliasis	1
Psychiatric Disorders	insomnia*	4
	[see Warnings and Precautions (5.6)]	
Nervous System Disorders	headache	6
	dizziness [see Warnings and Precautions (5.6)]	3
Respiratory, Thoracic and Mediastinal Disorders	dyspnea [see Warnings and Precautions (5.3)]	1
Gas trointes tinal Dis orders	nausea	7
	diarrhea	5
	constipation	3
	abdominal pain	2
	vomiting	2 2 2
	dyspepsia	2
Skin and Subcutaneous Tissue Disorders	rash [see Warnings and Precautions (5.3)]	2
	pruritus	
		1
Reproductive System and Breast Disorders	vaginitis	1^{\dagger}
	,	
General Disorders and Administration Site Conditions		1
	injection site reaction	1
**** #0#4	chest pain	1

^{*} N = 7274 † N = 3758 (women)

System/Organ Class	Adverse Reaction
Infections and Infestations	genital moniliasis
Blood and Lymphatic System Disorders	anemia thrombocytopenia
	granulocytopenia [see Warnings and Precautions (5.4)]
Immune System Disorders	allergic reaction [See Warnings and Precautions (5.3, 5.4)]
Metabolism and Nutrition Disorders	hyperglycemia hypoglycemia [see Warnings and Precautions (5.11)] hyperkalemia
Psychiatric Disorders	anxiety agitation confusion depression hallucination nightmare* [see Warnings and Precautions (5.6)] sleep disorder* anorexia abnormal dreaming*
Nervous System Disorders	tremor convulsions [see Warnings and Precautions (5.6)] paresthesia [see
	Warnings and Precautions (5.8)] vertigo hypertonia hyperkinesias abnormal gait somnolence* syncope
Respiratory, Thoracic and Mediastinal Disorders	epistaxis
Cardiac Disorders	cardiac arrest
	palpitation
	ventricular tachycardia
	ventricular arrhythmia
Vas cular Dis orders	phlebitis
Gas trointes tinal Dis orders	gastritis stomatitis pancreatitis esophagitis gastroenteritis glossitis pseudomembraneous/ C. difficile colitis [see Warnings and Precautions (5.7)]
Hepatobiliary Disorders	abnormal hepatic function
	increased hepatic enzymes
	increased alkaline phosphatase
Skin and Subcutaneous Tissue Disorders	urticaria [see Warnings and Precautions (5.3)]
Musculoskeletal and Connective Tissue Disorders	arthralgia tendinitis [see Warnings and Precautions (5.1)] myalgia skeletal pain
Renal and Urinary Disorders	abnormal renal function
	acute renal failure [see Warnings and Precautions (5.4)]
* N = 7274	

In clinical trials using multiple-dose therapy, ophthalmologic abnormalities, including cataracts and multiple punctate lenticular opacities, have been noted in patients undergoing treatment with quinolones, including levoltoxacin. The relationship of the drugs to these events is not presently established.

6.3 Postmarketing Experience

Table 6 lists adverse reactions that have been identified during post-approval use of levofloxacin. Because these reactions are reported voluntarily from a population of uncertain size, reliably estimating their frequency or establishing a causal relationship to drug exposure is not always possible.

Table 6: Postmarketing Reports Of Adverse Drug Reactions

System/Organ Class	Adverse Reaction
Blood and Lymphatic System Disorders	pancytopenia aplastic anemia leukopenia hemolytic anemia
Dioda and Lymphade System Distracts	[see Warnings and Precautions (5.4)] eosinophilia
Immune System Disorders	hypersensitivity reactions, sometimes fatal including:
,	anaphylactic/anaphylactoid reactions anaphylactic shock
	angioneurotic edema serum sickness
	[see Warnings and Precautions (5.3, 5.4)]
Psychiatric Disorders	psychosis paranoia isolated reports of suicide attempt and suicidal ideation
	[see Warnings and Precautions (5.6)]
Nervous System Disorders	exacerbation of myasthenia gravis [see Warnings and Precautions (5.2)]
	anosmia ageusia parosmia dysgeusia peripheral neuropathy [see Warnings
	and Precautions (5.8)] isolated reports of encephalopathy abnormal
	electroencephalogram (EEG) dysphonia pseudotumor cerebri [see Warnings
Eye Disorders	and Precautions (5.6)] vision disturbance, including diplopia
Eye Distruers	visual acuity reduced
	vision blurred
	scotoma
Ear and Labyrinth Disorders	hypoacusis
Dar and Eubyrman Distracts	tinnitus
Cardiac Disorders	isolated reports of torsade de pointes electrocardiogram OT prolonged
	[see Warnings and Precautions (5.9)] tachycardia
Vas cular Disorders	vasodilatation
Respiratory, Thoracic and Mediastinal Disorders	isolated reports of allergic pneumonitis [see Warnings and Precautions (5.4)]
Hepatobiliary Disorders	hepatic failure (including fatal cases) hepatitis jaundice
	[see Warnings and Precautions (5.4, 5.5)]
Skin and Subcutaneous Tissue Disorders	bullous eruptions to include: Stevens-Johnson Syndrome
	toxic epidermal necrolysis erythema multiforme
	[see Warnings and Precautions (5.4)] photosensitivity/phototoxicity reaction
	[see Warnings and Precautions (5.12)] leukocytoclastic vasculitis
Musculoskeletal and Connective Tissue Disorders	tendon rupture [see Warnings and Precautions (5.1)] muscle injury, including
	rupture rhabdomyolysis
Renal and Urinary Disorders	interstitial nephritis [see Warnings and Precautions (5.4)]
General Disorders and Administration Site Condition	
	pyrexia
Inves tig ations	prothrombin time prolonged
	international normalized ratio prolonged
	muscle enzymes increased

7. DRUG INTERACTIONS

7.1 Chelation Agents: Antacids, Sucralfate, Metal Cations, Multivitamins

LevofloxacinTablets

Levoltoxacm l'ablets
While the chelation by divalent cations is less marked than with other fluoroquinolones, concurrent
administration of levofloxacin tablets with antacids containing magnesium, or aluminum, as well as
sucralfate, metal cations such as iron, and multivitamin preparations with zinc may interfere with the
gastrointestinal absorption of levofloxacin, resulting in systemic levels considerably lower than
desired. Tablets with antacids containing magnesium, aluminum, as well as sucralfate, metal cations such
as iron, and multivitamins preparations with zinc or didanosine may substantially interfere with the
gastrointestinal absorption of levofloxacin, resulting in systemic levels considerably lower than
desired. These agents should be taken at least two hours before or two hours after oral levofloxacin
administration.

7.2 Warfarin

No significant effect of levofloxacin on the peak plasma concentrations, AUC, and other disposition parameters for R- and S- warfarin was detected in a clinical study involving healthy volunteers. Similarly, no apparent effect of warfarin on levofloxacin absorption and disposition was observed. However, there have been reports during the postmarketing experience in patients that levofloxacin enhances the effects of warfarin. Elevations of the prothrombin time in the setting of concurrent warfarin and levofloxacin use have been associated with episcodes of bleeding. Prothrombin time, International Normalized Ratio (INR), or other suitable anticoagulation tests should be closely monitored if levofloxacin is administered concomitantly with warfarin. Patients should also be monitored for evidence of bleeding [see Adverse Reactions (6.3); Patient Counseling Information (17.4)].

7.3 Antidiabetic Agents

Disturbances of blood glucose, including hyperglycenia and hypoglycenia, have been reported in patients treated concomitantly with fluoroquinolones and an antidiabetic agent. Therefore, careful monitoring of blood glucose is recommended when these agents are co-administered [see Warnings and Precautions (5.11); Adverse Reactions (6.2), Patient Counseling Information (17.4)].

7.4 Non-Steroidal Anti-Inflammatory Drugs

The concomitant administration of a non-steroidal anti-inflammatory drug with a fluoroquinolone, including levofloxacin, may increase the risk of CNS stimulation and convulsive seizures [see Warnings and Precautions (5.6)].

7.5 Theophylline

No significant effect of levofloxacin on the plasma concentrations, AUC, and other disposition parameters for theophylline was detected in a clinical study involving healthy volunteers. Similarly, no apparent effect of theophylline on levofloxacin absorption and disposition was observed. However, concomitant administration of other fluoroquinolones with theophylline has resulted in prolonged elimination half-life, elevated serum theophylline levels, and a subsequent increase in the risk of theophylline-related adverse reactions in the patient population. Therefore, theophylline levels should be closely monitored and appropriate dosage adjustments made when levofloxacin is co-administered. Adverse reactions, including seizures, may occur with or without an elevation in serum theophylline levels [see Warnings and Precautions (5.6)].

7.6 Cyclosporine

 $No\ significant\ effect\ of\ levo flox acin\ on\ the\ peak\ plasma\ concentrations,\ AUC,\ and\ other\ disposition$

parameters for cyclosporine was detected in a clinical study involving healthy volunteers. However, elevated serum levels of cyclosporine have been reported in the patient population when co-administered with some other fluoroquinolones. Levofloxacin C_{\max} and k_{∞} were slightly lower while T_{\max} and k_{δ} were slightly longer in the presence of cyclosporine than those observed in other studies without concomitant medication. The differences, however, are not considered to be clinically significant. Therefore, no dosage adjustment is required for levofloxacin or cyclosporine when administered concomitantly.

7.7 Digoxin

No significant effect of levofloxacin on the peak plasma concentrations, AUC, and other disposition parameters for digoxin was detected in a clinical study involving healthy volunteers. Levofloxacin absorption and disposition hierics were similar in the presence or absence of digoxin. Therefore, no dosage adjustment for levofloxacin or digoxin is required when administered concomitantly.

7.8 Probenecid and Cimetidine

No significant effect of probenecid or cimetidine on the C_{max} of levofloxacin was observed in a clinical study involving healthy volunteers. The AUC and b_0 of levofloxacin were higher while CL/F and CL_R were lower during concominant treatment of levofloxacin with probenecid or cimetidine compared to levofloxacin alone. However, these changes do not warrant dosage adjustment for levofloxacin when probenecid or cimetidine is co-administered.

7.9 Interactions with Laboratory or Diagnostic Testing

Some fluoroquinolones, including levofloxacin, may produce false-positive urine screening results for opiates using commercially available immunoassay kits. Confirmation of positive opiate screens by more specific methods may be necessary.

8. USE IN SPECIFIC POPULATIONS

8.1 Pregnancy

Pregnancy Category C. Levofloxacin was not teratogenic in rats at oral doses as high as 810 mg/kg/day which corresponds to 9.4 times the highest recommended human dose based upon relative body surface arra, or at infarous of Seesa sa high as 160 mg/kg/day corresponding to 1.9 times the highest state of the seesa shall be seed upon retained to the seesa shall be seed upon retained to the seesa seesa shall be seed upon retained to the seesa shall be seed upon retained between seesa shall be seed to the seesa shall be s when rabbits were dosed orally as high as 50 mg/kg/day which corresponds to 1.1 times the highest recommended human dose based upon relative body surface area, or when dosed intravenously as high as 25 mg/kg/day, corresponding to 0.5 times the highest recommended human dose based upon relative

There are, however, no adequate and well-controlled studies in pregnant women, Levofloxacin should be used during pregnancy only if the potential benefit justifies the potential risk to the fetus

8.3 Nursing Mothers

Based on data on other fluoroquinolones and very limited data on levofloxacin, it can be presumed that levofloxacin will be excreted in human milk. Because of the potential for serious adverse reactions from levofloxacin in nursing infants, a decision should be made whether to discontinue nursing or to discontinue the drug, taking into account the importance of the drug to the mother.

Quinolones, including levofloxacin, cause arthropathy and osteochondrosis in juvenile animals of several species. [see Warnings and Precautions (5.10) and Animal Toxicology and/or Pharmacology (13.2)].

Pharmacokinetics following intravenous administration

The pharmacokinetics of levofloxacin following a single intravenous dose were investigated in pediatric patients ranging in age from six months to 16 years. Pediatric patients cleared levofloxacin faster than adult patients resulting in lower plasma exposures than adults for a given mg/kg dose [see Clinical Pharmacology (12.3) and Clinical Studies (14.9)].

Inhalational Anthrax (Post-Exposure)

Levofloxacin is indicated in pediatric patients 6 months of age and older, for inhalational anthrax (post-exposure). The risk-benefit assessment indicates that administration of levofloxacin to pediatric patients is appropriate. The safety of levofloxacin in pediatric patients treated for more than 14 days has not been studied [see Indications and Usage (1.13), Dosage and Administration (2.2) and Clinical Studies (14.9)].

Plague

Levofloxacin is indicated in pediatric patients, 6 months of age and older, for treatment of plague, including pneumonic and septicentic plague due to Versinia pestis (Y. pestis) and prophylaxis for plague. Efficacy studies of levofloxacincould not be conducted in humans with pneumonic plague for ethical and feasibility reasons. Therefore, approval of this indication was based on an efficacy study conducted in animals. The risk-benefit assessment indicates that administration of levofloxacin to pediatric patients is appropriate [see Indications and Usage (1.14), Dosage and Administration (2.2) and Clinical Studies (14.10)].

Safety and effectiveness in pediatric patients below the age of six months have not been established.

In clinical trials, 1534 children (6 months to 16 years of age) were treated with oral and intraverous levofloxacin. Children 6 months to 5 years of age received levofloxacin 10 mg/kg wice a day and children greater than 5 years of age received 10 mg/kg once a day (maximum 500 mg per day) for approximately 10 days.

approximately to Joseph A subset of children in the clinical trials (1340 levofloxacin-treated and 893 non-fluoroquinolone-treated) enrolled in a prospective, long-term surveillance study to assess the incidence of protocol-defined musculoskeletal disorders (arthraigle, arthritis, tendinopathy, gait abnormality) during 60 days and 1 year following the first dose of study drug. Children treated with levofloxacin had a significantly higher incidence of musculoskeletal disorders when compared to the non-fluoroquinolone-treated children as illustrated in Table 7.

Table 7: Incidence of Musculos keletal Disorders in Pediatric Clinical Trial

Follow-up Period	Levofloxacin N = 1340	Non-Fluoroquinolone* N = 893	p-value†
60 days	28 (2.1%)	8 (0.9%)	p = 0.038
1 year‡	46 (3.4%)	16 (1.8%)	p = 0.025

Non-Fluoroquinolone: ceftriaxone, amoxicillin/ clavulanate, clarithromycin

Arthralgia was the most frequently occurring musculoskeletal disorder in both treatment groups. Most of the musculoskeletal disorders in both groups involved multiple weight-bearing joints. Disorders were moderate in 8/46 (17%) children and mild in 35/46 (76%) levofloxacin-treated children and most were treated with analgesics. The median time to resolution was 7 days for levofloxacin-treated children and 9 for non-fluoroquinolone-treated children (approximately 80% resolved within 2 months in both groups). No child had a severe or serious disorder and all musculoskeletal disorders resolved without sequelae.

Vomiting and diarrhea were the most frequently reported adverse events, occurring in similar frequency in the levofloxacin-treated and non-fluoroquinolone-treated children.

In addition to the events reported in pediatric patients in clinical trials, events reported in adults during clinical trials or post-marketing experience [see Adverse Reactions (6)] may also be expected to occur in pediatric patients.

as Geriatric use
Geriatric patients are at increased risk for developing severe tendon disorders including tendon rupture
when being treated with a fluoroquinolone such as levofloxacin. This risk is further increased in
patients receiving concomitant corticosteroid therapy. Tendinitis or tendon rupture can involve the
Achilles, hand, shoulder, or other tendon sites and can occur during or after completion of therapy;
cases occurring up to several months after fluoroquinolone treatment have been reported. Caution
should be used when prescribing levofloxacinto elderly patients especially those on corticosteroids.
Patients should be informed of this potential side effect and advised to discontinue levofloxacin and
contact their healthcare provider if any symptoms of tendinitis or tendon rupture occur [see Boxed
Warning; Warnings and Precautions (5.1); and Adverse Reactions (6.3)].

In phase 3 clinical trials, 1,945 levofloxacin-treated patients (26%) were \geq 65 years of age. Of these, 1,081 patients (14%) were between the ages of 65 and 74 and 864 patients (12%) were 75 years or older. No overall differences in safety or effectiveness were observed between these subjects and younger subjects, but greater sensitivity of some older individuals cannot be ruled out.

Severe, and sometimes fatal, cases of hepatotoxicity have been reported post-marketing in association with levofloxacin. The majority of fatal hepatotoxicity reports occurred in patients 65 years of age or older and most were not associated with hypersensitivity. Levofloxacinshould be discontinued immediately if the patient develops signs and symptoms of hepatitis [see Warnings and Precoutions (5.5)].

Elderly patients may be more susceptible to drug-associated effects on the QT interval. Therefore

^{†2-}sided Fisher's Exact Test

^{*}There were 1199 levofloxacin-treated and 804 non-fluoroguinolone-treated children who had a on visit. However, the incidence of musculoskeletal disorders was calculated using all reported events during the specified period for all children enrolled regardless of whether they completed the 1-year evaluation visit.

precaution should be taken when using levofloxacin with concomitant drugs that can result in prolongation of the QT interval (e.g., Class IA or Class III antiarrhythmics) or in patients with risk factors for torsade de pointes (e.g., known QT prolongation, uncorrected hypokalemia) [see Warnir and Precautions (5.9)].

The pharmacokinetic properties of levofloxacin in younger adults and elderly adults do not differ significantly when creatinine clearance is taken into consideration. However, since the drug is known to be substantially excreted by the kidney, the risks of toxic reactions to this drug may be greater in patients with impaired renal function. Because elderly patients are more likely to have decreased renal function, care should be taken in dose selection, and it may be useful to monitor renal function [see Clinical Pharmacology (12.3)].

8.6 Renal Impairment

Clearance of Evorlioxacin is substantially reduced and plasma elimination half-life is substantially prolonged in patients with impaired renal function (creatinine clearance < 50 mL/min), requiring dosage adjustment in such patients to avoid accumulation. Neither hemodialysis nor continuous ambulary peritoneal dialysis (CAPD) is effective in removal of levofloxacin from the body, indicating that supplemental doses of levofloxacin are not required following hemodialysis or CAPD [see Dosage and Administration (2.3)].

8.7 Hepatic Impairment

Pharmacokinetic studies in hepatically impaired patients have not been conducted. Due to the limited extent of levofloxacin metabolism, the pharmacokinetics of levofloxacin are not expected to be affected

In the event of an acute overdosage, the stomach should be emptied. The patient should be observed and appropriate hydration maintained. Levofloxacin is not efficiently removed by hemodialysis or peritoneal dialysis.

Levofloxacinexhibits a low potential for acute toxicity. Mice, rats, dogs and monkeys exhibited the following clinical signs after receiving a single high dose of levofloxacin ataxia, ptosis, decreasee locomotor activity, dyspnea, prostration, tremos, and comulsions. Doses in excess of 1500 mg/kg orally and 250 mg/kg IV produced significant mortality in rodents.

Levofloxacin is a synthetic broad-spectrum antibacterial agent for oral administration. Chemically, levofloxacin, a chiral fluorinated carboxyquimolone, is the pure (+)(5)-enantiomer of the racemic drug substance ofloxacin. The chemical amus is (+,(5))-ofluoro-2,-dihydyro-3-methyl-10-(4-methyl-1-piperazinyl)-7-oxo-7H-pyrido[1,2,3-de]-1,4-benzoxazine-6-carboxylic acid hemihydrate.

Figure 1: The Chemical Structure of Levofloxacin

The empirical formula is $C_{18}H_{20}FN_3O_4$ *½ H_2O and the molecular weight is 370.38. Levofloxacin is a white to light yellow crystalline powder. The molecule exists as a zwitterion at the pH conditions in the conditions of the conditions small intestine.

The data demonstrate that from pH 0.6 to 5.8, the solubility of levofloxacin is essentially constant (approximately 100 mg/mL). Levofloxacin is considered soluble to freely soluble in this pH range, as defined by USP nomenclature. Above pH 5.8, the solubility increases rapidly to its maximum at pH 6.7 (272 mg/mL) and is considered freely soluble in this range. Above pH 6.7, the solubility decreases and reaches a minimum value (about 50 mg/mL) at a pH of approximately 6.9.

Levofloxacin has the potential to form stable coordination compounds with many metal ions. This *in vitro* chelation potential has the following formation order: Al+3>Cu+2>Zn+2>Mg+2>Ca+2.

Excipients and Description of Dosage Forms

Levofloxacin Tablets

Levofloxacin tablets are available as film-coated tablets and contain the following inactive ingredients:

- Levolloxacin tablets are available as film-coated tablets and contain the following inactive ingredients:

 250 mg (as expressed in the anhydrous form): colloidal silicon dioxide, hypromellose, iron oxide red, magnesium stearate, microcrystalline cellulose, polyethylene glycol, polysorbate 80, povidone, propylene glycol, sodium starch glycolae and tiannium dioxide.

 500 mg (as expressed in the anhydrous form): colloidal silicon dioxide, hypromellose, iron oxide red, iron oxide yellow magnesium stearate, microcrystalline cellulose, polyethylene glycol, polysorbate 80, povidone, propylene glycol, sodium starch glycolae and tiannium dioxide.

 750 mg (as expressed in the anhydrous form): colloidal silicon dioxide, hypromellose, magnesium stearate, microcrystalline cellulose, polyethylene glycol, polysorbate 80, povidone, propylene glycol, sodium starch glycolate and tiannium dioxide.

12. CLINICAL PHARMACOLOGY

12.1 Mechanism of Action

Levofloxacin is a member of the fluoroquinolone class of antibacterial agents [see Microbiology (12.4)].

12.3 Pharmacokinetics

The mean ±SD pharmacokinetic parameters of levofloxacin determined under single and steady-state conditions following oral tablet, oral solution, or intravenous (IV) doses of levofloxacin are summarized in Table 8.

Table 8: Mean ±SD Levofloxacin PK Parameters

Regimen	Cmay (mcg/mL)	T _{may} (h)	AUC (mcg·h/mL)	CL/F1 (mL/min)	Vd/F2 (L)	t _{1/2} (h)	CL _R (mL/min)
Single dose							
250 mg oral tablet ³	2.8 ± 0.4	1.6 ± 1.0	27.2 ± 3.9	156 ± 20	ND	7.3 ± 0.9	142 ± 21
500 mg oral tablet ³ *	5.1 ± 0.8	1.3 ± 0.6	47.9 ± 6.8	178 ± 28	ND	6.3 ± 0.6	103 ± 30
500 mg oral solution ¹²	5.8 ± 1.8	0.8 ± 0.7	47.8 ± 10.8	183 ± 40	112 ± 37.2	7.0 ± 1.4	ND
500 mg IV ³	6.2 ± 1.0	1.0 ± 0.1	48.3 ± 5.4	175 ± 20	90 ± 11	6.4 ± 0.7	112 ± 25
750 mg oral tablet ⁵ *	9.3 ± 1.6	1.6 ± 0.8	101 ± 20	129 ± 24	83 ± 17	7.5 ± 0.9	ND
750 mg IV ⁵	11.5 ±4.0 ⁴	ND	110 ±40	126 ±39	75 ± 13	7.5 ± 1.6	ND
Multiple dose							
500 mg every 24h oral tablet ³	5.7 ± 1.4	1.1 ± 0.4	47.5 ± 6.7	175 ± 25	102 ± 22	7.6 ± 1.6	116 ± 31
500 mg every 24h IV ³	6.4 ± 0.8	ND	54.6 ± 11.1	158 ± 29	91 ± 12	7.0 ± 0.8	99 ± 28
500 mg or 250 mg every 24h IV, patients with bacterial infection ⁶		ND	72.5 ± 51.2^7	154 ± 72	111 ± 58	ND	ND
750 mg every 24h oral tablet ⁵	8.6 ± 1.9	1.4 ± 0.5	90.7 ± 17.6	143 ± 29		8.8 ± 1.5	116 ± 28
750 mg every 24h IV ⁵	12.1 ± 4.1^4	ND	108 ± 34	126 ± 37	80 ± 27	7.9 ± 1.9	ND
500 mg oral tablet single dose, effects of gender and age:							
Male ⁸	5.5 ± 1.1	1.2 ± 0.4	54.4 ± 18.9	166 ± 44		7.5 ± 2.1	126 ± 38
Female ⁹	7.0 ± 1.6	1.7 ± 0.5	67.7 ± 24.2	136 ± 44		6.1 ± 0.8	106 ± 40
Young 10	5.5 ± 1.0	1.5 ± 0.6	47.5 ± 9.8	182 ± 35	83 ± 18	6.0 ± 0.9	140 ± 33
Elderly ¹¹	7.0 ± 1.6	1.4 ± 0.5	74.7 ± 23.3	121 ± 33	67 ± 19	7.6 ± 2.0	91 ± 29
500 mg oral single dose tablet, patients with renal insufficience	y:						
CLCR 50 - 80 mL/min	7.5 ± 1.8	1.5 ± 0.5	95.6 ± 11.8	88 ± 10	ND	9.1 ± 0.9	57 ± 8
CLCR 20 - 49 mL/min	7.1 ± 3.1	2.1 ± 1.3	182.1 ± 62.6	51 ± 19	ND	27 ± 10	26 ± 13
CLCR <20 mL/min	8.2 ± 2.6	1.1 ± 1.0	263.5 ± 72.5	33 ± 8	ND	35 ± 5	13 ± 3
Hemodialysis	5.7 ± 1.0	2.8 ± 2.2	ND	ND	ND	76 ± 42	ND
CAPD	6.9 ± 2.3	1.4 ± 1.1	ND	ND	ND	51 ± 24	ND

- 1 clearance/bioavailability
- volume of distribution/bioavailability

- 2 volume of distribution/bioavalability
 3 healthy males 18-53 years of age
 4 60 min infusion for 250 mg and 500 mg doses, 90 min infusion for 750 mg dose
 5 healthy male and female subjects 18-54 years of age
 6 500 mg every 48h for patients with moderate renal impairment (CLCR 20-50 mL/min) and infections of the respiratory tract or skin
 7 dose-normalized values (to 500 mg dose), estimated by population pharmacokinetic modeling
 8 healthy females 18-80 years of age
 9 healthy females 18-80 years of age
 10 young healthy male and female subjects 18-36 years of age
 11 healthy deletrly male and female subjects 66-80 years of age
 12 healthy males and female subjects 66-80 years of age
 13 healthy males and female subjects 66-80 years of age
 14 healthy elderly male and female subjects 66-80 years of age
 15 healthy males and females 19-55 years of age.
 16 Absolute bioavalibility, F = 0.99 ± 0.08 from a 500 mg tablet and F = 0.99 ± 0.06 from a 750 mg tablet;
 18 ND=not determined.

Absorption

Levofloxacin is rapidly and essentially completely absorbed after oral administration. Peak plasma concentrations are usually attained one to two hours after oral dosing. The absolute bioavailability levofloxacin from a 500 mg tablet and a 750 mg tablet of levofloxacin are both approximately 99% demonstrating complete oral absorption of levofloxacin. Following a single intravenous dose of levofloxacin to healthy volunteers, the mean ±SD peak plasma concentration attained was 6.2 ± 1.0 mg/ml. after a 500 mg dose infused over 60 minutes and 11.5 ± 4.0 mg/ml. after a 750 mg dose infused over 60 minutes and 11.5 ± 4.0 mg/ml. after a 750 mg dose infused over 90 minutes.

infused over 90 minutes. Levofloxacin Oral Solution and Tablet formulations are bioequivalent. Levofloxacin pharmacokinetics are linear and predictable after single and multiple oral or IV dosing regimers. Steady-state conditions are reached within 48 hours following a 500 mg or 750 mg once-daily dosage regimen. The mean ±5D peak and trough plasma concentrations attained following multiple once-daily oral dosage regimens were approximately 5.7 ± 1.4 and 0.5 ± 0.2 mg/ml. after the 500 mg doses, and 8.6 ±1.9 and 1.1 ± 0.4 mcg/ml. after the 750 mg doses, respectively. The mean ± SD peak and trough plasma concentrations attained following multiple once-daily IV regimens were approximately 6.4 ± 0.8 and 0.5 ± 0.2 mcg/ml. after the 500 mg doses, and 1.1 ± 4.1 and 1.3 ± 0.71 mcg/ml. after the 750 mg doses, respectively. Oral administration of a 500 mg dose of levofloxacin with food prolongs the time to neak concentration by a proxyximately 1.0 hour and decreases the neak concentration by the time to peak concentration by approximately 1 hour and decreases the peak concentration by approximately 25% following called and approximately 14% following tablet and approximately 25% following oral solution administration. Therefore, levofloxacin tablets can be administered without regard to food.

The plasma concentration profile of levofloxacin after IV administration is similar and comparable in The photocock of the province of the province

Figure 2: Mean Levofloxacin Plasma Concentration vs. Time Profile: 750 mg

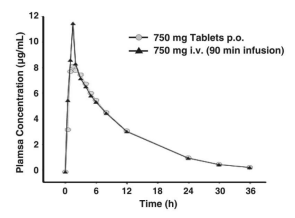
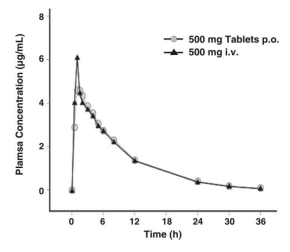


Figure 3: Mean Levofloxacin Plasma Concentration vs. Time Profile: 500 mg



The mean volume of distribution of levofloxacin generally ranges from 74 to 112 L after single and multiple 500 mg or 750 mg dosses, indicating widespread distribution into body tissues. Levofloxacin reaches its peak levels in skin tissues and in blister fluid of healthy subjects at approximately 3 hours after dosing. The skin tissue biopsy to plasma AUC ratio is approximately 2 and the blister fluid to plasma AUC ratio is approximately 1 following multiple once-daily oral administration of 750 mg and 500 mg dosses of levofloxacin, respectively, no healthy subjects. Levofloxacin also penetrates well into lung tissues. Lung tissue concentrations were generally 2- to 5-fold higher than plasma concentrations and ranged from approximately 2.4 to 11.3 mc/g over a 24-hour period after a single 500 mg oral dose.

In vitro, over a clinically relevant range (1 to 10 mcg/mL) of serum/plasma levofloxacin concentrations, levofloxacin is approximately 24 to 38% bound to serum proteins across all species studied, as determined by the equilibrium dialysis method. Levofloxacin is mainly bound to serum albumin in humans. Levofloxacin binding to serum proteins is independent of the drug concentration.

Levofloxacin is stereochemically stable in plasma and urine and does not invert metabolically to its enantiomer, D-ofloxacin. Levofloxacin undergoes limited metabolism in humans and is primarily excreted as unchanged drug in the urine. Following oral administration, approximately 87% of an administered dose was recovered as unchanged drug in urine within 48 hours, whereas less than 4% of the dose was recovered in feces in 72 hours. Less than 5% of an administered dose was recovered in the urine as the desmethyl and N-oxide metabolites, the only metabolites identified in humans. These metabolites have little relevant pharmacological activity.

Excretion

Levofloxacin is excreted largely as unchanged drug in the urine. The mean terminal plasma elimination half-life of levofloxacin ranges from approximately 6 to 8 hours following single or multiple doses of levofloxacin given-orally or intraverously. The mean apparent total body clearance and renal clearance range from approximately 144 to 226 mL/min and 95 to 142 mL/min, respectively. Renal clearance in excess of the glomerular filtration rate suggests that tubular secretion of levofloxacin occurs in addition to its glomerular filtration. Concornitar administration of either cimetidine or probenecid results in approximately 24% and 35% reduction in the levofloxacin renal clearance, respectively, indicating that secretion of levofloxacin occurs in the renal proximal tubules. No levofloxacin crystals were found in any of the urine samples freshly collected from subjects receiving levofloxacin.

There are no significant differences in levofloxacin pharmacokinetics between young and elderly subjects when the subjects' differences in creatinine clearance are taken into consideration. Following a 500 mg oral dose of levofloxacin to healthy elderly subjects (66-80 years of age), the mean terminal plasma elimination half-life of levofloxacin was about 7.6 hours, as compared to approximately 6 hours

in younger adults. The difference was attributable to the variation in renal function status of the subjects and was not believed to be clinically significant. Drug absorption appears to be unaffected by age. Levofloxaci

Pediatrics

The pharmacokinetics of levofloxacin following a single 7 mg/kg intravenous dose were investigated in pediatric patients ranging in age from 6 months to 16 years. Pediatric patients cleared levofloxacin faster than adult patients, resulting in lower plasma exposures than adults for a given mg/kg dose. Subsequent pharmacokinetic analyses predicted that a dosage regimen of 8 mg/kg every 12 hours (not to exceed 250 mg per dose) for pediatric patients 6 months to 17 years of age would achieve comparable steady state plasma exposures (AUC₀₋₂₄ and C_{max}) to those observed in adult patients administered 500 mg of levofloxacin once every 24 hours.

There are no significant differences in levofloxacin pharmacokinetics between male and female subjects when subjects' differences in creatinine clearance are taken into consideration. Following a 500 mg oral dose of levofloxacin to healthy male subjects, the mean terminal plasma elimination half-life of levofloxacin was about 7.5 hours, as compared to approximately 6.1 hours in female subjects. This difference was attributable to the variation in renal function status of the male and female subjects and was not believed to be clinically significant. Drug absorption appears to be unaffected by the gender of the subjects. Dose adjustment based on gender alone is not necessary.

The effect of race on levofloxacin pharmacokinetics was examined through a covariate analysis performed on data from 72 subjects: 48 white and 24 non-white. The apparent total body clearance and apparent volume of distribution were not affected by the race of the subjects.

Clearance of levofloxacin is substantially reduced and plasma elimination half-life is substantially prolonged in adult patients with impaired renal function (creatinine clearance < 50 mL/min, requiring dosage adjustment in such patients to avoid accumulation. Neither hemodialysis nor continuous ambulatory peritoneal dialysis (CAPD) is effective in removal of levofloxacin from the body, indicating that supplemental doses of levofloxacin are not required following hemodialysis or CAPD [see Dosage and Administration (2.3), Use in Specific Populations (8.6)].

Hepatic Impairment

Pharmacokinetic studies in hepatically impaired patients have not been conducted. Due to the limited extent of levofloxacin metabolism, the pharmacokinetics of levofloxacin are not expected to be affected by hepatic impairment [See Use in Specific Populations (8.7)].

Bacterial Infection

The pharmacokinetics of levofloxacin in patients with serious community-acquired bacterial infections are comparable to those observed in healthy subjects.

Drug-Drug Interactions

The potential for pharmacokinetic drug interactions between levofloxacin and antacids warfarin, theophylline, cyclosporine, digoxin, probenecid, and cimetidine has been evaluated [see Drug Interactions (7)].

12.4 Microbiology

Mechanism of Action

Levofloxacin is the L-isomer of the racemate, ofloxacin, a quinolone antimicrobial agent. The antibacterial activity of ofloxacin resides primarily in the L-isomer. The mechanism of action of levofloxacin and other fluoroquinolone antimicrobials involves inhibition of bacterial topoisomerase IV and DNA gyrase (both of which are type II topoisomerases), enzymes required for DNA replication, transcription, repair and recombination.

Mechanism of Resistance

Fluoroquinolone resistance can arise through mutations in defined regions of DNA gyrase or topoisomerase IV, termed the Quinolone-Resistance Determining Regions (QRDRs), or through altered efflux.

Fluoroquinolones, including levofloxacin, differ in chemical structure and mode of action from aminoglycosides, macrolides and β -lactam antibiotics, including penicillins. Fluoroquinolones may, therefore, be active against bacteria resistant to these antimicrobials.

Resistance to levofloxacin due to spontaneous mutation *in vitro* is a rare occurrence (range: 10^{-9} to 10^{-10}). Cross-resistance has been observed between levofloxacin and some other fluoroquinolones, some microorganisms resistant to other fluoroquinolones may be susceptible to levofloxacin.

Activity in vitro and in vivo

 $Levo flox a cin \ has \ \textit{in vitro} \ activity \ against \ Gram-negative \ and \ Gram-positive \ bacteria.$

 $Levo flox actin has been shown to be active against most isolates of the following bacteria both {\it in vitro} and in clinical infections as described in {\it Indications and Usage (1):}$

Gram-Positive Bacteria

Enterococcus faecalis

Staphylococcus aureus (methicillin-susceptible isolates)

Staphylococcus epidermidis (methicillin-susceptible isolates)

Staphylococcus saprophyticus

Streptococcus pneumoniae (including multi-drug resistant isolates [MDRSP])¹

Streptococcus pyogenes

1 MDRSP (Multi-drug resistant Streptococcus pneumoniae) isolates are isolates resistant to two or more of the following antibiotics: penicillin (MIC ≥2 mcg/mL), 2nd generation cephalosporins, e.g., cefuroxime; macrolides, tetracyclines and trimethoprim/sulfamethoxazole.

Gram-Negative Bacteria

Enterobacter cloacae

Escherichia coli

Haemophilus influenzae

Haemophilus parainfluenzae Klebsiella pneumoniae

Legionella pneumophila

Moraxella catarrhalis

Proteus mirabilis

Pseudomonas aeruainosa

Serratia marcescens

Other Bacteria Chlamydophila pneumoniae

Mycoplasma pneumoniae

The following in vitro data are available, but their clinical significance is unknown:

 $Levo flox a cin exhibits \textit{ in vitro } minimum inhibitory concentrations (MIC values) of 2 \,mcg/mL or less$ against most (290%) isolates of the following microorganisms; however, the safety and effectiveness of levofloxacin in treating clinical infections due to these bacteria have not been established in adequate and well-controlled clinical trials.

Gram-Positive Bacteria

Staphylococcus haemolyticus

β-hemolytic Streptococcus (Group C/F)

β-hemolytic Streptococcus (Group G)

Streptococcus agalactiae

Streptococcus milleri

Viridans group streptococci

Bacillus anthraci

Gram-Negative Bacteria

Acinetobacter baumannii Acinetohacter Iwoffii

Bordetella pertussis

Citrobacter koseri Citrobacter freundii

Enterobacter aerogenes

Enterobacter sakazakii

Klebsiella oxytoca

Morganella morganii

Pantoea agglomerans

Proteus vulaaris

Providencia rettaeri Providencia stuartii

Pseudomonas fluorescens

Yersinia pestis

Anaerobic Gram-Positive Bacteria

Clostridium perfringens

Susceptibility Tests

When available, the clinical microbiology laboratory should provide the results of *in vitro* susceptibility test results for antimicrobial drug products used in the 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 MIC values should be determined using a standardized procedure. Standardized procedures are based on a dilution method-2²⁴ (froth or agapt or equivalent with standardized inoculum concentrations and standardized concentrations of levofloxacin powder. The MIC values should be interpreted according to the criteria outlined in Table 9.

Diffusion techniques:

Quantitative methods that require measurement of zone diameters also provide reproducible estimates Quantum emittibility of bacteria to antimicrobial compounds. One such standardized procedure^{2,3} of the susceptibility of bacteria to antimicrobial compounds. One such standardized procedure^{2,3} requires the use of standardized inoculum concentrations. This procedure uses paper disks impregnated with 5 mtg levofloxacin to exhibit the susceptibility of bacteria to levofloxacin.

Reports from the laboratory providing results of the standard single-disk susceptibility test with a 5 mcg levofloxacin disk should be interpreted according to the criteria outlined in Table 9.

Table 9: Sus ceptibility Test Interpretive Criteria for Levofloxacin

	Minimum Inhibitory Co	ncentratio	ns (mcg/mL)	Disk Diffu	sion (zone diame	ter in mm)
Pathogen	S	I	R	S	I	R
Enterobacteriaceae	≤2	4	≥8	≥17	14-16	≤13
Enterococcus faecalis	≤2	4	≥8	≥17	14-16	≤13
Staphylococcus species	≤2	4	≥8	≥17	14-16	≤13
Pseudomonas aeruginosa	≤2	4	≥8	≥17	14-16	≤13
Haemophilus influenzae	≤2	_†	-	≥17	-	
Haemophilus parainfluenzae	≤2	-	-	≥17	-	
Streptococcus pneumoniae	≤2	4	≥8	≥17	14-16	≤13
Streptococcus pyogenes	≤2	4	≥8	≥17	14-16	≤13
Yersinia pestis ⁴	≤0.25	-	-		-	
Bacillus anthracis ⁴	≤0.25	-	-	-	-	-

A report of Susceptible indicates that the pathogen is likely to be inhibited if the antimicrobial compound in the blood reaches the concentrations usually achievable. 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 which prevents small uncontrolled technical factors from causing major discrepancies in interpretation. A report of Resistant indicates that the pathogen is not tilely to be inhibited if the antimicrobial compound in the blood reaches the concentrations usually achievable; 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 individuals performing the test. ^{1,2,3,4} Standard levofloxacin powder should provide the range of MIC values moted in Table 10. For the diffusion technique using the 5 mcg disk, the criteria in Table 10 should be achieved.

Table 10: Quality Control for Susceptibility Testing

Microorganism	Microorg-anism QC Number	MIC (mcg/mL)	Disk Diffusion (zone diameter in mm)
Enterococcus faecalis	ATCC 29212	0.25 - 2	
Escherichia coli	ATCC 25922	0.008 - 0.06	29 – 37
Escherichia coli	ATCC 35218	0.015 - 0.06	
Haemophilus influenzae	ATCC 49247	0.008 - 0.03	32 – 40
Pseudomonas aeruginosa	ATCC 27853	0.5 – 4	19 – 26
Staphylococcus aureus	ATCC 29213	0.06 - 0.5	
Staphylococcus aureus	ATCC 25923		25 – 30
Streptococcus pneumoniae	ATCC 49619	0.5 - 2	20 – 25

13. NONCLINICAL TOXICOLOGY

13.1 Carcinogenesis, Mutagenesis, Impairment Of Fertility

13.1 Carcmogenesis, Mutagenesis, Impairment OI Ferthity
I al lifetime bioassay in rats, levofloxacin exhibited no carcinogenic potential following daily dietary
administration for 2 years; the highest dose (100 mg/kg/day) was 1.4 times the highest recommended
human dose (750 mg) based upon relative body surface area. Levofloxacin did not shorten the time to
tumor development of UV-induced skin tumors in hairless allino (Skh-1) mice at any levofloxacin dose
level and was therefore not photo-carcinogenic under conditions of this study. Dermal levofloxacin
concentrations in the hairless mice ranged from 25 to 42 mc/g at the highest levofloxacin dose level
(300 mg/kg/day) used in the photo-carcinogenicity study. By comparison, dermal levofloxacin
concentrations in human subjects receiving 750 mg of levofloxacin averaged approximately 11.8 mcg/g at C_{max}.

Levofloxacin was not mutagenic in the following assays: Ames bacterial mutation assay (S. typhimurium and E. coli), CHO/HGPRT forward mutation assay, mouse micronucleus test, mouse dominant lethal test, rat unscheduled DNA synthesis assay, and the mouse sister chromatid exchange assay. It was positive in the in vitro chromosomal aberration (CHL cell line) and sister chromatid exchange (CHL/IU cell line)

Levo floxacin caused no impairment of fertility or reproductive performance in rats at oral doses as high as 360 mg/kg/day, corresponding to 4.2 times the highest recommended human dose based upon relative body surface area and intravenous doses as high as 100 mg/kg/day, corresponding to 1.2 times the highest recommended human dose based upon relative body surface area.

13.2 Animal Toxicology And/Or Pharmacology

Levofloxacin and other quinolones have been shown to cause arthropathy in immature animals of most Levofloxacin and other quinolones have been shown to cause arthropathy in immature animals of most species tested [see Warmings and Precoutions (5.10]). In immature dogs (4-5 months old), or all osses of 10 mg/kg/day for 7 days and intravenous doses of 40 mg/kg/day for 14 days of levofloxacin resulted in arthropathic lesions. Administration at oral doses of 300 mg/kg/day for 7 days and intravenous doses of 60 mg/kg/day for 4 weeks produced arthropathy in juvenile rats. Three-month old beagle dogs dosed orally with levofloxacin at 40 mg/kg/day exhibited clinically severe arthrotoxicity resulting in the termination of dosing at Day 8 of a 14-day dosing routine. Slight musculoskeleal clinical effects, in the absence of gross pathological or histopathological effects, resulted from the lowest dose level of 2.5 mg/kg/day (approximately 0.2-fold the pediatric dose based upon AUC comparisons). Sprovist and articular cartilage lesions were observed at the 10 and 40 mg/kg/dose levels (approximately 0.7-fold and 2.4-fold the pediatric dose, respectively, based on AUC comparisons). Articular cartilage gross pathology and histopathology persisted to the end of the 18-week recovery period for those dogs from the 10 and 40 mg/kg/day dose levels.

When tested in a mouse ear swelling bioassay, levofloxacin exhibited phototoxicity similar in magnitude to ofloxacin, but less phototoxicity than other quinolones.

While crystalluria has been observed in some intravenous rat studies, urinary crystals are not formed in the bladder, being present only after micturition and are not associated with nephrotoxicity.

In mice, the CNS stimulatory effect of quinolones is enhanced by concomitant administration of non-

steroidal anti-inflammatory drugs

t absence of data on resistant isolates precludes defining any categories other than "Susceptible." Isolates yiel diameter results suggestive of a "nonsusceptible" category should be submitted to a reference laboratory for

In dogs, levofloxacin administered at 6 mg/kg or higher by rapid intravenous injection produced hypotensive effects. These effects were considered to be related to histamine release.

In vitro and in vivo studies in animals indicate that levofloxacin is neither an enzyme inducer nor inhibitor in the human therapeutic plasma concentration range; therefore, no drug metabolizing enzyme-related interactions with other drugs or agents are anticipated.

14.1 Nosocomial Pneumonia

Adult patients with clinically and radiologically documented nosocomial pneumonia were enrolled in a multicenter, randomized, open-label study comparing intravenous levofloxacin (750 mg once daily) for lotal of 7-15 days to intravenous imipenemcilastatin (500-1000 mg every 6-8 hours daily) for alotal of 7-15 days to intravenous imipenemchiastatin (500-1000 mg every 6-8 hours daily) followed by oral ciprofloxacin (750 mg every 12 hours daily) for a total of 7-15 days. Levofloxacin-treated patients received an average of 7 days of intravenous therapy (range: 1-16 days); comparator-treated patients received an average of 8 days of intravenous therapy (range: 1-19 days).

days of intravenous therapy (range: 1-19 days). Overall, in the clinically and microbiologically evaluable population, adjunctive therapy was empirically initiated at study entry in 56 of 93 (60.2%) patients in the levofloxacin arm and 53 of 94 (36.4%) patients in the comparator arm. The average duration of adjunctive therapy was 7 days in the levofloxacin arm and 7 days in the comparator. In clinically and microbiologically evaluable patients with documented Pseudomonas aeruginosa infection, 15 of 17 (88.2%) received ceftazidine (N=11) or piperacillilirazioabactam (N=4) in the levofloxacin arm and 16 of 17 (94.1%) received an animoglycoside in the comparator arm. Overall, in clinically and microbiologically evaluable patients, vancomycin was added to the treatment regimen of 37 of 93 (39.8%) patients in the levofloxacin arm and 28 of 94 (29.8%) patients in the comparator arm for suspected methicillin-resistants. S. aureus infection.

Clinical success rates in clinically and microbiologically evaluable patients at the posttherapy visit (primary study endpoint assessed on day 3-15 after completing therapy) were 58.1% for levofloxacin and 60.6% for comparator. The 95% CI for the difference of response rates (levofloxacin minus comparator) was [1-72, 12.0]. The microbiological eradication rates at the posttherapy visit were 66.7% for levofloxacin and 60.6% for comparator. The 95% CI for the difference of eradication rates (levofloxacin minus comparator) was [-8.3, 2 0.3]. Clinical success and microbiological eradication rates by pathogen are detailed in Table 11.

Table 11: Clinical Success Rates and Bacteriological Eradication Rates (Nosocomial Pneumonia)

	_	1	_	
Pathogen	N	Levofloxacin No. (%) of Patients Microbiologic/ Clinical Outcomes	Ν	Imipenem/Cilastatin No. (%) of Patients Microbiologic/ Clinical Outcomes
MSSA*	21	14 (66.7)/13 (61.9)	19	13 (68.4)/15 (78.9)
P. aeruginosa [†]	17	10 (58.8)/11 (64.7)	17	5 (29.4)/7 (41.2)
S. marcescens	11	9 (81.8)/7 (63.6)	7	2 (28.6)/3 (42.9)
E. coli	12	10 (83.3)/7 (58.3)	11	7 (63.6)/8 (72.7)
K. pneumoniae‡	11	9 (81.8)/5 (45.5)	7	6 (85.7)/3 (42.9)
H. influenzae	16	13 (81.3)/10 (62.5)	15	14 (93.3)/11 (73.3)
S. pneumoniae	4	3 (75.0)/3 (75.0)	7	5 (71.4)/4 (57.1)

14.2 Community-Acquired Pneumonia: 7-14 day Treatment Regimen

14.2 Community-Acquired Pneumonia: 7-14 day Treatment Regimen

Adult inpatients and outpatients with a diagnosis of community-acquired bacterial pneumonia were
evaluated in 2 pivotal clinical studies. In the first study, 590 patients were emolled in a prospective,
multi-center, unblinded randomized trial comparing levofloxacin 500 mg once daily orally or
intravenously once or in equally divided
doses twice daily followed by cefuroxime axetil 500 mg orally twice daily for a total of 7 to 14 days.
Patients assigned to treatment with the control regimen were allowed to receive erythromycin (or
doxycycline if intolerant of erythromycin) if an infection due to atypical pathogers was suspected or
proven. Clinical and microbiologic evaluations were performed during treatment, 5 to 7 days
posttherapy, and 3 to 4 weeks posttherapy, Clinical success (cure plus improvement) with levofloxacin
at 5 to 7 days posttherapy, the primary efficacy variable in this study, was superior (95%) to the control
group (83%). The 59% Cf for the difference of response rates (levofloxacin minus comparator) was [-6,19]. In the second study, 264 patients were enrolled in a prospective, multi-center, non-comparative
trial of 500 mg levofloxacin administered orally or intravenously once daily for 7 to 14 days. Clinical
success for clinically evaluable patients was 93%. For both studies, the clinical success rate in patients
with atypical pneumonia due to Chlamydophila pneumoniae, and Legionella
pneumophila were 96%, 96%, and 70%, respectively. Microbiologic eradication rates across both

Table 12: Bacteriological Eradication Rates Across 2 Community Acquired Pneumonia Clinical Studies

Pathogen	No. Pathogens	Bacteriological Eradication Rate (%)
H. influenzae	55	98
S. pneumoniae	83	95
S. aureus	17	88
M. catarrhalis	18	94
H. parainfluenzae	19	95
K. pneumoniae	10	100

Community-Acquired Pneumonia Due to Multi-Drug Resistant Streptococcus pneumonia

Levofloxacin was effective for the treatment of community-acquired pneumonia caused by multi-drug resistant Streptococcus pneumoniae (MDRSP). MDRSP isolates are isolates resistant to two or more of the following amtibacterials: pencillilin (MCZ 2 mcg/ml), 2md generation cephalosporins (e.g., cefuroxime, macrolides, tetracyclines and trinethoprim/sulfamethoxazole). Of 40 microbiologically evaluable patients with MDRSP isolates, 38 patients (950-98) achieved clinical and bacteriologic success at post-therapy. The clinical and bacterial success rates are shown in Table 13.

Table 13: Clinical and Bacterial Success Rates for Levofloxacin-Treated MDRSP in Community Acquired Pneumonia Patients (Population Valid for Efficacy)

Screening Susceptibility	Clinical Success		Bacteriological Success*	
	n/N [†]	%	n/N [‡]	%
Penicillin-resistant	16/17	94.1	16/17	94.1
2 nd generation Cephalosporin resistant	31/32	96.9	31/32	96.9
Macrolide-resistant	28/29	96.6	28/29	96.6
Trimethoprim/ Sulfamethoxazole resistant	17/19	89.5	17/19	89.5
Tetracycline-resistant	12/12	100	12/12	100

One patient had a respiratory isolate that was resistant to tetracycline, cefuroxime, macrolides and TMP/SMX and One patient had a respiratory solute that was resistant to terracyctine, ceturoxime, macrobies and IMP/SMA and intermediate to pencillian and a blood solute that was intermediate to pencillian and a blood solute that was intermediate to pencillian and reduroxime and resistant to the other classes. The patient is included in the database based on respiratory isolute.

†=n=the number of microbiologically evaluable patients who were clinical successes; N=number of microbiologically evaluable patients in the designated resistance group.

†=n=the number of MDRSP isolates eradicated or presumed eradicated in microbiologically evaluable patients; N=number of MDRSP isolates in a designated resistance group.

Not all isolates were resistant to all antimicrobial classes tested. Success and eradication rates are summarized in Table 14.

Table 14: Clinical Success and Bacteriologic Eradication Rates for Resistant Streptococcus pneumoniae (Community Acquired Pneumonia)

Type of Resistance	Clinical Success	Bacteriologic Eradication
Resistant to 2 antibacterials	17/18 (94.4%)	17/18 (94.4%)
Resistant to 3 antibacterials	14/15 (93.3%)	14/15 (93.3%)
Resistant to 4 antibacterials	7/7 (100%)	7/7 (100%)
Resistant to 5 antibacterials	0	0
Bacteremia with MDRSP	8/9 (89%)	8/9 (89%)

14.3 Community-Acquired Pneumonia: 5-day Treatment Regimen

tay to 10 days.

Clinical success rates (cure plus improvement) in the clinically evaluable population were 90.9% in the levofloxacin 750 mg group and 91.1% in the levofloxacin 500 mg group. The 95% CI for the difference of response rates (levofloxacin 750 minus levofloxacin 500) was [-5.9, 5.4]. In the clinically evaluable population (31-38 days after enrollment) pneumonia was observed in 7 out of 151 patients in the levofloxacin 750 mg group and 2 out of 147 patients in the levofloxacin 500 mg group. Given the small numbers observed, the significance of this finding cannot be determined statistically. The microbiological efficacy of the 5-day regimen was documented for infections listed in Table 15.

[|] S. pneumonine | 4 | 3 (73.0) | 1 / 1 | 2 - 2 - 3 (73.0) | 1 / 1 | 2 - 2 - 3 (73.0) | 3 (73.0) | 1 / 1 | 2 - 2 - 3 (73.0) | 3 (73.0) | 3 (73.0) | 3 (73.0) | 4 | 3 (73.0) | 5 (73.0) | 5 (73.0) | 5 (73.0) | 5 (73.0) | 5 (73.0) | 5 (73.0) | 5 (73.0) | 5 (73.0) | 5 (73.0) | 5 (73.0) | 5 (73.0) | 5 (73.0) | 5 (73.0) | 5 (73.0) | 5 (73.0) | 5 (73.0) | 5 (73.0) | 5 (73.0) | 5 (73.0) | 5 (73.0) | 5 (73.0) | 5 (73.0) | 5 (73.0) | 5 (73.0) | 5 (73.0) | 5 (73.0) | 5 (73.0) | 5 (73.0) | 5 (73.0) | 5 (73.0) | 5 (73.0) | 5 (73.0) | 5 (73.0) | 5 (73.0) | 5 (73.0) | 5 (73.0) | 5 (73.0) | 5 (73.0) | 5 (73.0) | 5 (73.0) | 5 (73.0) | 5 (73.0) | 5 (73.0) | 5 (73.0) | 5 (73.0) | 5 (73.0) | 5 (73.0) | 5 (73.0) | 5 (73.0) | 5 (73.0) | 5 (73.0) | 5 (73.0) | 5 (73.0) | 5 (73.0) | 5 (73.0) | 5 (73.0) | 5 (73.0) | 5 (73.0) | 5 (73.0) | 5 (73.0) | 5 (73.0) | 5 (73.0) | 5 (73.0) | 5 (73.0) | 5 (73.0) | 5 (73.0) | 5 (73.0) | 5 (73.0) | 5 (73.0) | 5 (73.0) | 5 (73.0) | 5 (73.0) | 5 (73.0) | 5 (73.0) | 5 (73.0) | 5 (73.0) | 5 (73.0) | 5 (73.0) | 5 (73.0) | 5 (73.0) | 5 (73.0) | 5 (73.0) | 5 (73.0) | 5 (73.0) | 5 (73.0) | 5 (73.0) | 5 (73.0) | 5 (73.0) | 5 (73.0) | 5 (73.0) | 5 (73.0) | 5 (73.0) | 5 (73.0) | 5 (73.0) | 5 (73.0) | 5 (73.0) | 5 (73.0) | 5 (73.0) | 5 (73.0) | 5 (73.0) | 5 (73.0) | 5 (73.0) | 5 (73.0) | 5 (73.0) | 5 (73.0) | 5 (73.0) | 5 (73.0) | 5 (73.0) | 5 (73.0) | 5 (73.0) | 5 (73.0) | 5 (73.0) | 5 (73.0) | 5 (73.0) | 5 (73.0) | 5 (73.0) | 5 (73.0) | 5 (73.0) | 5 (73.0) | 5 (73.0) | 5 (73.0) | 5 (73.0) | 5 (73.0) | 5 (73.0) | 5 (73.0) | 5 (73.0) | 5 (73.0) | 5 (73.0) | 5 (73.0) | 5 (73.0) | 5 (73.0) | 5 (73.0) | 5 (73.0) | 5 (73.0) | 5 (73.0) | 5 (73.0) | 5 (73.0) | 5 (73.0) | 5 (73.0) | 5 (73.0) | 5 (73.0) | 5 (73.0) | 5 (73.0) | 5 (73.0) | 5 (73.0) | 5 (73.0) | 5 (73.0) | 5 (73.0) | 5 (73.0) | 5 (73.0) | 5 (73.0) | 5 (73.0) | 5 (73.0) | 5 (73.0) | 5 (73.0) | 5 (73.0) | 5 (73.0) | 5 (73.0) | 5 (73.0) | 5 (73.0) | 5 (73.0) | 5 (73.0) | 5 (73.0) | 5 (73.0) | 5 (73.0) | 5 (73.0) | 5 (73.0) | 5 (73.0) | 5 (73.0) | 5 (73.0) | 5 (

Table 15: Bacteriological Eradication Rates (Community-Acquired Pneumonia)

S. pneumoniae	19/20 (95%)
Haemophilus influenzae	12/12 (100%)
Haemophilus parainfluenzae	10/10 (100%)
Mycoplasma pneumoniae	26/27 (96%)
Chlamydophila pneumoniae	13/15 (87%)

14.4 Acute Bacterial Sinusitis: 5-day and 10-14 day Treatment Regimens

Levofloxacin is approved for the treatment of acute bacterial sinusitis (ABS) using either 750 mg by mouth x5 days or 500 mg by mouth once daily x 10-14 days. To evaluate the safety and efficacy of a high dose short course of levofloxacin, 780 outpatient adults with clinically and radiologically determined acute bacterial sinusitis were evaluated in a double-blind, randomized, prospective, multicenter study comparing levofloxacin 750 mg by mouth once daily for five days to levofloxacin 500 mg by mouth once daily for 10 days.

Clinical success rates (defined as complete or partial resolution of the pre-treatment signs and symptoms of ABS to such an extent that no further antibiotic treatment was deemed necessary) in the microbiologically evaluable population were 91.4% (139/152) in the levofloxacin 750 mg group and 88.6% (132/149) in the levofloxacin 500 mg group at the test-of-cure (TOC) visit (95% CI [-4.2, 10.0] for levofloxacin 750 mg minus levofloxacin 500 mg).

Rates of clinical success by pathogen in the microbiologically evaluable population who had specimens obtained by artral tap at study entry showed comparable results for the five- and ter regimens at the test-of-cure visit 22 days post treatment.

Table 16: Clinical Success Rate by Pathogen at the TOC in Microbiologically Evaluable Subjects Who Underwent Antral Puncture (Acute Bacterial Sinusitis)

Pathogen	Levofloxacin 750 mg x 5 days	Levofloxacin 500 mg x 10 days
Streptococcus pneumoniae*	25/27 (92.6%)	26/27 (96.3%)
Haemophilus influenzae*	19/21 (90.5%)	25/27 (92.6%)
Moraxella catarrhalis*	10/11 (90.9%)	13/13 (100%)

^{*} Note: Forty percent of the subjects in this trial had specimens obtained by sinus endoscopy. The efficacy data for subjects whose specimen was obtained endoscopically were comparable to those presented in the above table

14.5 Complicated Skin and Skin Structure Infections

14.5 Complicated Skin and Skin Structure Intections
Three hundred ninety-nine patients were enrolled in an open-label, randomized, comparative study for complicated skin and skin structure infections. The patients were randomized to receive either levofloxacin 750 mg once daily (IV followed by oral), or an approved comparator for a median of 10 ± 4.7 days. As is expected in complicated skin and skin structure infections, surgical procedures were performed in the levofloxacin and comparator groups. Surgery (incision and drainage or debridement) was performed on 45% of the levofloxacin-treated patients and 44% of the comparator-treated patients, either shortly before or during antibiotic treatment and formed an integral part of therapy for this indication.

Among those who could be evaluated clinically 2-5 days after completion of study drug, overall success rates (improved or cured) were 116/138 (84.1%) for patients treated with levofloxacin and 106/132 (80.3%) for patients treated with the comparator.

Success rates varied with the type of diagnosis ranging from 68% in patients with infected ulcers to 90% in patients with infected wounds and abscesses. These rates were equivalent to those seen with comparator drugs.

14.6 Chronic Bacterial Prostatitis

Adult patiers with a clinical diagnosis of prostatitis and microbiological culture results from urine sample collected after prostatic massage (VB₃) or expressed prostatic secretion (EPS) specimens obtained via the Meares-Stamey procedure were enrolled in a multicenter, randomized, double-blind study comparing oral leverolloxacin 500 mg, once daily for a total of 28 days to oral ciprofloxacin 500 mg, twice daily for a total of 28 days. The primary efficacy endpoint was microbiologic efficacy in microbiologically evaluable patients. A total of 136 and 125 incrobiologically evaluable patients were enrolled in the levofloxacin and ciprofloxacin groups, respectively. The microbiologic eradication rate by patient infection at 5-18 days after completion of therapy was 75.0% in the levofloxacin group (95% CI-1-2.58, 8.38) for levofloxacin minus ciprofloxacin). The overall eradication rates for pathogens of interest are presented in Table 17.

Table 17: Bacteriological Eradication Rates (Chronic Bacterial Prostatitis)

Levofloxacin (N=136)		Cip	rofloxacin (N=125)	
Pathogen	N	Eradication	N	Eradication
E. coli	15	14 (93.3%)	11	9 (81.8%)
E. faecalis	54	39 (72.2%)	44	33 (75.0%)
C anidarmidic*	11	0 (01 00/)	1.4	11 (70 60/)

b. epidermidis* | 11 | 9 (81.8%) | 14 | 11 (78.6%) * Eradication rates shown are for patients who had a sole pathogen only; mixed cultures were excluded.

Eradication rates for S. epidermidis when found with other co-pathogens are consistent with rates seen in pure isolates.

Clinical success (cure + improvement with no need for further antibiotic therapy) rates in climical success (use "improvement with a freet or further attanced that are in microbiologically evaluable population 5-18 days after completion of therapy were 75.0% for levofloxacin-treated patients and 72.8% for ciprofloxacin-treated patients (95% CI [-8.87, 13.27] for levofloxacin minus ciprofloxacin). Clinical long-term success (24-45 days after completion of therapy) rates were 66.7% for the levofloxacin-treated patients and 76.9% for the ciprofloxacin-treated patients (95% CI [-23.40, 2.89] for levofloxacin minus ciprofloxacin).

14.7 Complicated Urinary Tract Infections and Acute Pyelonephritis: 5-day Treatment Regimen

14.7 Complicated Urinary Tract Infections and Acute Pyelonephritis: 5-day Treatment Regimen To evaluate the safety and efficacy of the higher dose and shorter course of levofloxacin, 1109 patients with cUT1 and AP were enrolled in a randomized, double-blind, multicenter clinical trial conducted in the US from November 2004 to April 2006 comparing levofloxacin 750 mg IV or orally once daily for 5 days (364 patients) with ciprofloxacin 400 mg IV or 500 mg orally wice daily for 10 days (653 patients). Patients with AP complicated by underlying renal diseases or conditions such as complete obstruction, surgery, transplanation, concurrent infection or congenial mulformation were excluded. Efficacy was measured by bacteriologic eradication of the baseline organism(s) at the post-therapy visit in patients with a pathogen identified at baseline. The post-therapy (test-of-cure) visit occurred 10 to 14 days after the last active dose of levofloxacin and 5 to 9 days after the last dose of active citorafloxacin. active ciprofloxacin.

The bacteriologic cure rates overall for levofloxacin and control at the test-of-cure (TOC) visit for the group of all patients with a documented pathogen at baseline (modified intent to treat or mITT) and the group of patients in the mITT population who closely followed the protocol (Microbiologically Evaluable) are summarized in Table 18.

Table 18: Bacteriological Eradication at Test-of-Cure

	Levofloxacin 750 mg orally or IV once daily for 5 days		Ciprofloxacin 400 mg IV/500 mg orally twice daily for 10 days		Overall Difference [95% CI]
	n/N	%	n/N	%	Levofloxacin- Ciprofloxacin
	mITT Population*				
Overall (cUTI or AP)	252/333	75.7	239/318	75.2	0.5 (-6.1, 7.1)
cUTI	168/230	73.0	157/213	73.7	
AP	84/103	81.6	82/105	78.1	
	Mi	crobiologically	Evaluable Population [†]		
Overall (cUTI or AP)	228/265	86.0	215/241	89.2	-3.2 [-8.9, 2.5]
cUTI	154/185	83.2	144/165	87.3	
AP	74/80	92.5	71/76	93.4	

[AP 74/80 y 32.5]. Thr mITT population included patients who received study medication and who had a positive (=10⁵CFU/mL) urise culture with no more than 2 uropathogens at baseline. Patients with missing response were counted as failures in this analysis.

The Microbiologically Evaluable population included patients with a confirmed diagnosis of cUTI or AP, a causative organism(s) at baseline present at = 10⁵ CFU/mL, a valid test-of-cure urine culture, no pathogen isolated from blood resistant to study drug, no premature discontinuation or loss to follow-up, and compliance with reatment (among other criteria).

 $\label{thm:microbiologic} Microbiologically\ Evaluable\ population\ at\ TOC\ for\ individual\ pathogens\ recovered\ from\ patients\ randomized\ to\ levofloxacin\ treatment\ are\ presented\ in\ Table\ 19.$

Table 19: Bacteriological Eradication Rates for Individual Pathogens Recovered From Patients Randomized to Levofloxacin 750 mg QD for 5 Days Treatment

Pathogen	Bacteriological Eradication Rate (n/N)	%
Escherichia coli*	155/172	90
Klebsiella pneumoniae	20/23	87
Proteus mirabilis	12/12	100

*The predominant organism isolated from patients with AP was *E. coli*: 91% (63/69) eradication in AP and 89% (92/103) in patients with eUTI.

To evaluate the safety and efficacy of the 250 mg dose, 10 day regimen of levofloxacin, 567 patients with uncomplicated UTI, mild-to-moderate cUTI, and mild-to-moderate AP were enrolled in a randomized, double-blind, multicenter clinical trial conducted in the US from June 1993 to January 1995 comparing levofloxacin 250 orally once daily for 10 days (285 patients) with ciprofloxacin 500 mg orally twice daily for 10 days (282 patients). Patients with a resistant pathogen, recurrent UTI, women over age 55 years, and with an indwelling catheter were initially excluded, prior to protocol amendment which took place after 30% of emollment. Microbiological efficacy was measured by bacteriologic eradication of the baseline organism(s) at 1-12 days post-therapy in patients with a pathogen identified at baseline.

The bacteriologic cure rates overall for levofloxacin and control at the test-of-cure (TOC) visit for the group of all patients with a documented pathogen at baseline (modified intent to treat or mTTT) and the group of patients in the mTTT population who closely followed the protocol (Microbiologically Evaluable) are summarized in Table 20.

Table 20. Bacteriological Eradication Overall (cUTI or AP) at Test-Of-Cure

	Levofloxacin		Ciprofloxacin		
	250 mg once daily for 10 days		500 mg twice daily for 10 days		
	n/N %		n/N	%	
mITT Population [†]	174/209	83.3	184/219	84.0	
Microbiologically	164/177	92.7	159/171	93.0	
Evaluable Population [‡]					

^{* 1-9} days posttherapy for 30% of subjects enrolled prior to a protocol amendment; 5-12 days posttherapy for 709

14.3 mindational Attuirax (Vest-Exposure)
The effectiveness of levolfoxacin for this indication is based on plasma concentrations achieved in humans, a surrogate endpoint reasonably likely to predict clinical benefit. Levofloxacin has not been tested in humans for the post-exposure prevention of inhalation atthax. The mean plasma concentration of levofloxacin associated with a statistically significant improvement in survival over placebo in the rhesus monkey model of inhalational anthrax are reached or exceeded in adult and pediatric patients receiving the recommended oral and intravenous dosage regimens [see Indications and Usage (1.13); Dosage and Administration (2.1, 2.2)].

 $Levo flox a cin pharmacokinetics \ have \ been \ evaluated \ in \ a dult \ and \ pediatric \ patients. \ The \ mean \ (\pm \ SD)$ Levotroxacin paarmaconaetics and seener valuated in adult and peduaric patients. In the mean $(\pm 5D)$ steady state peak plasma concentration in human adults receiving 500 mg orally or intravenously once daily is 5.7 ± 1.4 and 6.4 ± 0.8 mcg/ml., respectively; and the corresponding total plasma exposure $(AUC_{p,2d})$ is $AY \pm 6.7$ and 5.4 ± 11.1 mcg, hulm, respectively. The predicted steady-state pharmacokinetic parameters in pediatric patients ranging in age from 6 months to 17 years receiving 8 mg/kg orally every 12 hours (not to exceed 250 mg per doss) were calculated to be comparable to those observed in adults receiving 500 mg orally once daily [see Clinical Pharmacology (12.3)].

In adults, the safety of levofloxacin for treatment durations of up to 28 days is well characterized. However, information pertaining to extended use at 500 mg daily up to 60 days is limited. Prolonged levofloxacin therapy in adults should only be used when the benefit outweighs the risk.

In pediatric patients, the safety of levofloxacin for treatment durations of more than 14 days has not been in pecuairt patients, are safety or territoristat flow the deadurin quantities of inflict and it supplies studied. An increased incidence of musculoskeletal adverse evens (arthraligia, arthritis, tendinopathy, gait abnormality) compared to control shas been observed in clinical studies with treatment duration of up to 14 days. Long-term safety data, including effects on cartilage, following the administration of levofloxacin to pedularite patients is limited [see Warnings and Precautions (5.10), Use in Specific Populations (8.41).

A placebo-controlled animal study in rhesus monkeys exposed to an inhaled mean dose of 49 $\rm LD_{50}$ (-2.7 \times 10°) spores (range 17-118 $\rm LD_{50}$) of B. anthracis (Ames strain) was conducted. The minimbilitory concentration (MIC) of levofloxacin for the anthrax strain used in this study was 0.125 inhibitory concentration (MLC) of tevoltoxacin for the anthrax strain used in this study was 0.125 meg/mL. In the animals studied, mean plasma concentrations of levolfoxacin achieved at expected T_{max} (I hour post-dose) following oral dosing to steady state ranged from 0.279 to 4.87 mcg/mL. Steady state trough concentrations at 24 hours post-dose ranged from 0.107 to 0.164 mcg/mL. Mean (SD) steady state AUC_{0.24} was 33.4 ± 3.2 mcg.h/mL (range 30.4 to 36.0 mcg.h/mL). Mortality due to anthrax for animals that received a 30 day regimen of oral levolfoxacin beginning 24 hrs post exposure was significantly lower (1/10), compared to the placebog group (9/10) [P=0.0011, 2-sided Fisher's Exact Test]. The one levolfoxacin treated animal that died of anthrax did so following the 30-day drug administration period.

14.10 Plague

Efficacy studies of levofloxacin could not be conducted in humans with pneumonic plague for ethical and feasibility reasons. Therefore, approval of this indication was based on an efficacy study conducted the study of the st

The mean plasma concentrations of levofloxacin associated with a statistically significant improvement in survival over placebo in an African green monkey model of pneumonic plague are reached or exceeded in adult and pediatric patients receiving the recommended oral and intravenous dosage regimens [see Indications and Usage (1.14), Dosage and Administration (2.1), (2.2)].

Levofloxacin pharmacokinetics has been evaluated in adult and pediatric patients. The mean (\pm SD) steady state peak plasma concentration in human adults receiving 500 mg orally or intravenously once daily is 5.7 ± 1.4 and 6.4 ± 0.8 mcg/mL, respectively; and the corresponding total plasma exposure (AUC_{0.24}) is 47.5 ± 6.7 and 5.4 ± 11.1 mcg, h/mL, respectively. The predicted steady-state pharmacokinetic parameters in pediatric patients ranging in age from 6 months to 17 years receiving 8 mg/kg orally every 12 hours (not to exceed 250 mg per dose) were calculated to be comparable to those observed in adults receiving 500 mg orally once daily [see Clinical Pharmacology (12.3)].

observed in adults receiving 500 mg orally once daily [see Clinical Pharmacology (12.3)]. A placebo-controlled animal study in African green monkeys exposed to an inhaled mean dose of 65 LD₅₀ (range 3 to 145 LD₅₀) of Yersinia pestis (CO92 strain) was conducted. The minimal inhibitory concentration (MIC) of levofloxacin for the Y. pestis strain used in this study was 0.03 mcg/ml. Mean plasma concentrations of levofloxacin achieved at the end of a single 30min infusion ranged from 2.84 to 3.50 mcg/ml. in African green monkeys. Trough concentrations at 24 hours post-dose ranged from color: blue; was 11.9 (3.1) mcg, brinl. (range 9.50 to 16.86 mcg, h/ml.). Animals were randomized to receive either a 10-day regimen of i.v. levofloxacin or placebo beginning within 6 hrs of the onset of telemetered fever (2-39°C for more than 1 hour). Mortality in the levofloxacin group was significantly lower (117) compared to the placebo group (77) [p-0.001, Fisher's Exact Test, exact 95% confidence interval (-99.9%, -55.5%) for the difference in mortality]. One levofloxacin-treated animal was euthanized on Day 9 post-exposure to Y. pestis due to a gastric complication; it had a blood culture positive for Y. pestis on Day 3 and all subsequent daily blood cultures from Day 4 through Day 7 were negative.

- 1. Clinical and Laboratory Standards Institute (CLSI). Methods for Dilution Antimicrobial Susceptibility Tests for Bacteria That Grow Aerobically, Approved Standard – 9th ed. CLSI Document M7-A9, CLSI, 950 West Valley Rd., Suite 2500, Wayne, PA, 2012.

 2. CLSI. Performance Standards for Admirctobial Susceptibility Testing; 22nd Informational Supplement. CLSI Document M100 – S22, 2012.
- CLSI Performance Standards for Antimicrobial Disk Susceptibility Tests. Approved Standard 11th ed. CLSI M2-A11, 2012.
- et. CLSI Me2-K11, 2012.

 C.LSI Methods for Antimicrobial Dilution and Disk Susceptibility Testing of Infrequently Isolated or Fastidious Bacteria; Approved Guideline 2nd ed. CLSI Document M45-A2, 2010.

16. HOW SUPPLIED/STORAGE AND HANDLING

16.1 Levofloyacin Tablets

Levofloxacin Tablets are supplied as 250, 500, and 750 mg capsule-shaped, coated tablets. Levofloxacin Tablets are packaged in bottles and in unit-dose blister strips in the following configurations:

- 250 mg tablets are pink and are debossed; "W" on one side and "544" on the other side

- 250 mg tablets are plink and are debos:
 bottle of 20 (NDC 64679-544-01)
 bottle of 50 (NDC 64679-544-02)
 bottle of 100 (NDC 64679-544-03)
 bottle of 500 (NDC 64679-544-04)
- unit-dose packages of 100 (NDC 64679-544-05)
- 500 mg tablets are peach and are debossed: "W" on one side and "545" on the other side
 bottle of 20 (NDC 64679-545-01)
 bottle of 50 (NDC 64679-545-02)

- bottle of 100 (NDC 64679-545-03)
- bottle of 500 (NDC 64679-545-04) unit-dose packages of 100 (NDC 64679-545-05)
- 750 mg tablets are white and are debossed: "W" on one side and "547" on the other side
 bottle of 20 (NDC 64679-547-01)
 bottle of 50 (NDC 64679-547-02)

- bottle of 100 (NDC 64679-547-03)
 bottle of 500 (NDC 64679-547-04)

[†]The mITT population included patients who had a pathogen isolated at baseline. Patients with missing response ed as failures in this analysis.

^{*}The Microbiologically Evaluable population included mITT patients who met protocol-specified evaluability criteria.

• unit-dose packages of 100 (NDC 64679-547-05)

Store at 20°-25°C (68°-77°F) [See USP Controlled Room Temperature]. Keep the container closed

17. PATIENT COUNSELING INFORMATION

See FDA-Approved Medication Guide (17.6)

17.1 Antibacterial Resistance

Antibacterial drugs including levofloxacin should only be used to treat bacterial infections. They do not treat viral infections (e.g., the common cold). When levofloxacin 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 levofloxacin or other antibacterial drugs in the future. drugs in the future.

17.2 Administration with Food, Fluids, and Concomitant Medications

Patients should be informed that levofloxacin tablets may be taken with or without food. The tablet should be taken at the same time each day.

Patients should drink fluids liberally while taking levofloxacin to avoid formation of a highly concentrated urine and crystal formation in the urine.

Antacids containing magnesium, or aluminum, as well as sucralfate, metal cations such as iron, and multivitamin preparations with zinc or didanosine should be taken at least two hours before or two hours after oral levofloxacin administration.

17.3 Serious and Potentially Serious Adverse Reactions

- Patients should be informed of the following serious adverse reactions that have been associated with levofloxacin or other fluoroquinolone use:

 Tendon Dis orders: Patients should contact their healthcare provider if they experience pain, swelling, or inflammation of a tendon, or weakness or inability to use one of their joints; rest and refrain from exercise; and discontinue levofloxacin treatment. The risk of severe tendon disorders with fluoroquinolones is higher in older patients usually over 60 years of age, in patients taking corticosteroid drugs, and in patients usually over 60 years of age, in patients taking corticosteroid drugs, and in patients usually over 60 years of age, in patients taking corticosteroid drugs, and in patients this thickney, heart or lung transplants.

 Exacerbation of Myas thenia Gravis: Patients should inform their physician of any history of myasthenia gravis. Patients should be informed that levofloxacin can cause hypersensitivity reactions: Patients should be informed that levofloxacin can cause hypersensitivity reactions. even following the first dose. Patients should discontinue the drug at the first sign of a skin rash, hives or other skin reactions, a rapid heartheat, difficulty in swallowing or breathing, any swelling suggesting angioederme (e.g., swelling of the lips, tongue, face, tightness of the throat, hoarseness), or other symptoms of an allergic reaction.

 Hepatotoxicity: Severe hepatotoxicity (including acute hepatitis and fatal events) has been reported in patients taking levofloxacin. Patients should inform their physician and be instructed to discontinue levofloxacin treatment immediately if they experience any signs or symptoms of liver injury including: loss of appetite, nausea, vomiting, fever, weakness, tiredness, right upper quadrant tenderness, tiching, yellowing of the skin and eyes, light colored bowel movements or dark colored urine.
- Convulsions: Convulsions have been reported in patients taking fluoroquinolones, including levofloxacin. Patients should notify their physician before taking this drug if they have a history of
- convulsions.

 Neurologic Adverse Effects (e.g., dizziness, lightheadedness, increased intracranial pressure): Patients should know how they react to levofloxacin before they operate an automobile or machinery or engage in other activities requiring mental alertness and coordination. Patients should notify their physician if persisters headache with or without blurred vision occurs.

 Diarrhea: 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.
- Peripheral Neuropathies: If symptoms of peripheral neuropathy including pain, burning, tingling, numbness, and/or weakness develop, patients should discontinue treatment and contact their physician.
- Prolongation of the QT Interval: Patients should inform their physician of any personal or family history of QT prolongation or proarrhythmic conditions such as hypokalemia, bradycardia, or recent myocardial ischemia; if they are taking any Class IA (quindine, procainamide), or Class III (amiodarone, soalola) antiarrhythmic agents. Patients should notify their physicians if they have any symptoms of prolongation of the QT interval, including prolonged heart palpitations or a loss of
- consciousness.

 * Musculos keletal Disorders in Pediatric Patients: Parents should inform their child's physician if their child has a history of joint-related problems before taking this drug. Parents of pediatric patients should also notify their child's physician of any tendon or joint-related problems that occur during or following levofloxacin therapy Isee Warnings and Precautions (3.10) and Use in Specific Populations (8.41).

 **Photosensitivity/Phototoxicity: Patients should be advised that photosensitivity/phototoxicity has been reported in patients receiving fluoroquinolone antibiotics. Patients should minimize or avoid exposure to natural or artificial smilght (taming beds or UVAB Teatmenty while taking fluoroquinolones, If patients need to be outdoors when taking fluoroquinolones, they should wear loose-fitting clothes that protect shaft from sun exposure and discuss other sun protection measures with their physician. If a sunburn like reaction or skin eruption occurs, patients should contact their physician.

17.4 Drug Interactions with Insulin, Oral Hypoglycemic Agents, and Warfarin

Patients should be informed that if they are diabetic and are being treated with insulin or an oral hypoglycemic agent and a hypoglycemic reaction occurs, they should discontinue levofloxacin and consult a physician.

Patients should be informed that concurrent administration of warfarin and levofloxacin has been associated with increases of the International Normalized Ratio (INR) or prothrombin time and clinical episodes of bleeding. Patients should notify their physician if they are taking warfarin, be monitored for evidence of bleeding, and also have their anticoagulation tests closely monitored while taking warfarin

Patients given levofloxacin tablets for these conditions should be informed that efficacy studies could not be conducted in humans for ethical and feasibility reasons. Therefore, approval for these conditions was based on efficacy studies conducted in animals.

Manufactured by:

Wockhardt Limited,

Mumbai, India. Distributed by

Wockhardt USA LLC

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Parsippany, NJ 07054

USA.

Rev.171012

17.6 FDA-Approved Medication Guide

MEDICATION GUIDE

LEVOFLOXACIN TABLETS

250 mg, 500 mg and 750 mg

Read this Medication Guide before you start taking levofloxacin and each time you get a refill. There may be new information. This Medication Guide does not take the place of talking to your healthcare provider about your medical condition or your treatment.

What is the most important information I should know about levofloxacin tablets? Levofloxacin, a fluoroquinolone antibiotic, can cause serious side effects. Some of these serious side effects

If you have any of the following serious side effects while you take levofloxacin tablets, get medical help right away. Talk with your healthcare provider about whether you should continue to take levofloxacin tablets.

1. Tendon rupture or swelling of the tendon (tendinitis)

Tendon problems can happen in people of all ages who take levofloxacin tablets. Tendons are tough cords of tissue that connect muscles to bones.

Some tendon problems include pain, swelling, tears, and inflammation of tendons including the back of

the ankle (Achilles), shoulder, hand, or other tendon sites

- The risk of getting tendon problems while you take levofloxacin tablets are higher if you:
- o are over 60 years of age
- o are taking steroids (corticosteroids)
- o have had a kidney, heart, or lung transplant,
- Tendon problems can happen in people who do not have the above risk factors when they take levofloxacin tablets
- · Other reasons that can increase your risk of tendon problems can include:
- o physical activity or exercise
- o kidney failure
- o tendon problems in the past, such as in people with rheumatoid arthritis (RA).
- Call your healthcare provider right away at the first sign of tendon pain, swelling or inflammation. Stop taking levofloxacin tablets until tendinitis or tendon rupture has been ruled out by your healthcare provider. Avoid exercise and using the affected area.

The most common area of pain and swelling is the Achilles tendon at the back of your ankle. This can also happen with other tendons. Talk to your healthcare provider about the risk of tendon rupture with continued use of levofloxacin tablets. You may need a different antibiotic that is not a fluoroquinolone to treat your infection.

- Tendon rupture can happen while you are taking or after you have finished taking levofloxacin tablets. Tendon ruptures have happened up to several months after people have finished taking their fluoroquinolone.
- Get medical help right away if you get any of the following signs or symptoms of a tendon rupture:
- o hear or feel a snap or pop in a tendon area
- o bruising right after an injury in a tendon area
- o unable to move the affected area or bear weight
- 2. Worsening of myasthenia gravis (a problem that causes muscle weakness). Fluoroquinolones like levofloxacin tablets may cause worsening of myasthenia gravis symptoms, including muscle weakness and breathing problems. Call your healthcare provider right away if you have any worsening muscle weakness or breathing problems.

• See "What are the possible side effects of levofloxacin tablets?"

What is Levofloxacin?

Levofloxacin is a fluoroquinolone antibiotic medicine used in adults age 18 years or older to treat certain infections caused by certain germs called bacteria. These bacterial infections include:

- nosocomial pneumonia
- community-acquired pneumonia acute sinus infection

- acute worsening of chronic bronchitis skin infections, complicated and uncomplicated chronic prostate infection urinary tract infections, complicated and uncomplicated

because plague and anthrax could not be studied in people.

- acute kidney infection (pyelonephritis)
- inhalational anthrax
- plague

Studies of levofloxacin tablets for use in the treatment of plague and anthrax were done in animals only.

Levofloxacin tablets are also used to treat children who are 6 months of age or older and may have breathed in anthrax germs, have plague, or been exposed to plague germs

It is not known if levofloxacin tablets are safe and effective in children under 6 months of age.

The safety and effectiveness in children treated with levofloxacin tablets for more than 14 days is not

Who should not take levofloxacin tablets?

Do not take levofloxacin tablets if you have ever had a severe allergic reaction to an antibiotic known as a fluoroquinolone, or if you are allergic to levofloxacin or any of the ingredients in levofloxacin tablets. See the end of this leaflet for a complete list of ingredients in levofloxacin tablets.

What should I tell my healthcare provider before taking levofloxacin tablets?

Before you take levofloxacin tablets, tell your healthcare provide if you:

- have tendon problems
- have a problem that causes muscle weakness (myasthenia gravis)
 have central nervous system problems such as seizures (epilepsy)

- Inave rever problems
 have nerve problems
 have or anyone in your family has an irregular heartbeat, especially a condition called "QT prolongation"
 have low blood potassium (hypokalemia)

- have bone problems have joint problems including rheumatoid arthritis (RA)
- have kidney problems. You may need a lower dose of levofloxacin tablets if your kidneys do not work well.

- work well.

 have liver problems

 have diabetes or problems with low blood sugar (hypoglycemia)

 are pregnant or plan to become pregnant. It is not known if levofloxacin will harm your unborn child.

 are breastfeeding or plan to breastfeed. It is not known if levofloxacin passes into your breast milk.

 You and your healthcare provider should decide if you will take levofloxacin tablets or breastfeed. You should not do both.

Tell your healthcare provider about all the medicines you take, including prescription and non-prescription medicines, vitamins, and herbal supplements.

Levofloxacin tablets and other medicines can affect each other causing side effects.

- Especially tell your healthcare provider if you take:
- a steroid medicine. An anti-psychotic medicine
- A tricyclic antidepressant
- A water pill (ultretic) certain medicines may keep levofloxacin tablets from working correctly. Take levofloxacin tablets either 2 hours be fore or 2 hours after taking these medicines or supplements:
- \boldsymbol{o} an antacid, multivitamin, or other medicines or supplements that have magnesium, aluminum, iron, or zinc
- o sucralfate (Carafate®)
- didanosine (Videx®.Videx® EC)

- o didanosine (Videx®, Videx® EC)
 a ablood thinner (warfain, Coumadin, Jantoven)
 a noral anti-diabetes medicine or insulin
 an NSAID (Mon-Steroidal Anti-Inflammatory Drug), Many common medicines for pain relief are
 NSAIDs. Taking an NSAID while you take levofloxacin tablets or other fluoroquinolones may
 increase your risk of certain hervous system effects and seizures.

 theophylline (Theo-24®, Elixophyllin®, Theochron®, Uniphyl®, Theolair®)
- · a medicine to control your heart rate or rhythm (antiarrhythmics)

Ask your healthcare provider if you are not sure if any of your medicines are listed above.

Know the medicines you take. Keep a list of your medicines and show it to your healthcare provider and pharmacist when you get a new medicine.

How should I take levofloxacin tablets?

- Trues should I take revoluntacin tablets?

 Take levofloxacin tablets exactly as your healthcare provider tells you to take it.

 Take levofloxacin tablets at about the same time each day.

 Drink pleny of fluids while you take levofloxacin tablets.

 Levofloxacin tablets can be taken with or without food.

 If you miss a does of levofloxacin tablets, take it as soon as you remember. Do not take more than 1 dose in 1 day.
- O no tskip any doses of levofloxacin tablets or stop taking it, even if you begin to feel better, until you finish your prescribed treatment unless:

o you have tendon problems. See "What is the most important information I should know about levofloxacin tablets?".

o $\,$ you have a serious allergic reaction. See "What are the possible side effects of levofloxacin tablets?".

o your healthcare provider tells you to stop taking levofloxacin tablets

levofloxacin tablets, it may mean that the bacteria causing your infection may be resistant to levofloxacin tablets. If your infection does not get better, call your healthcare provider. If your infection does not get better, levofloxacin tablets and other similar antibiotic medicines may not work for you in the future.

• If you take too much levofloxacin tablets, call your healthcare provider or get medical help right

- What should I avoid while taking levofloxacin tablets?

 Levofloxacin tablets can make you feel dizzy and lightheaded. Do not drive, operate machinery, or do other activities that require mental alertness or coordination until you know how levofloxacin
- 4. Other activities that require mema activities to contaminate and the sun. Levofloxacin tablets received with a contamining beds, and try to limit your time in the sun. Levofloxacin tablets can make your skin sersitive to the sun (photosensitivity) and the light from sunlamps and tanning beds. You could get severe surburn, blisters or swelling of your skin. If you get any of these symptoms while taking levofloxacin tablets, call your healthcare provider right away. You should use a sunscreen and wear a hat and clothes that cover your skin if you have to be in sunlight.

What are the possible side effects of levofloxacin tablets?

- Levofloxacin tablets can cause serious side effects, including:

 See "What is the most important information I should know about levofloxacin tablets?"
- · Serious allergic reactions

Allergic reactions can happen in people taking fluoroquinolones, including levofloxacin, even after only I dose. Stop taking levofloxacin tablets and get emergency medical help right away if you get any of the following symptoms of a severe allergic reaction:

- o hives
- o trouble breathing or swallowing
- o swelling of the lips, tongue, face
- o throat tightness, hoarseness
- o rapid heartbeat

Skin rash may happen in people taking levofloxacin tablets, even after only 1 dose. Stop taking levofloxacin tablets at the first sign of a skin rash and call your healthcare provider. Skin rash may be a sign of a more serious reaction to levofloxacin tablets.

Liver damage (hepatotoxicity): Hepatotoxicity can happen in people who take levofloxacin tablets.
 Call your healthcare provider right away if you have unexplained symptoms such as:

0	nausea or vomiting	o unusual tiredness
0	stomach pain	o loss of appetite
0	fever	o light colored bowel
0	weakness	movements
0	abdominal pain or tenderness	o dark colored urine
		o yellowing of your skin or
0	itching	the whites of your eyes

Stop taking levofloxacin tablets and tell your healthcare provider right away if you have yellowing of your skin or white part of your eyes, or if you have dark urine. These can be signs of a serious reaction to levofloxacin tablets (a liver problem).

Central Nervous System Effects. Seizures have been reported in people who take fluoroquinolone antibiotics including levofloxacin tablets. Tell your healthcare provider if you have a history of seizures. Ask your healthcare provider whether taking levofloxacin tablets will change your risk of having a seizure. Central Nervous System (CNS) side effects may happen as soon as after taking the first dose of levofloxacin tablets. Talk to your healthcare provider right away if you get any of these side effects, or other changes in mood or behavior:

o seizures	o trouble sleeping
o hear voices, see there	o nightmares
things, or sense	o feel lightheaded
things that are not	o feel more suspicious
(hallucinations)	(paranoia)
o feel restless	o suicidal thoughts or acts
o tremors	o a headache that will not
o feel anxious or nervous	go away, with or without blurred vision.
o confusion	
o depression	

• Intestine infection (Pseudomembranous colitis)

Pseudomembranous colitis can happen with many artibiotics, including levofloxacin tablets. Call your healthcare provider right away if you get watery diarrhea, diarrhea that does not go away, or bloody stools. You may have stomach cramps and a fever. Pseudomembranous colitis can happen 2 or more months after you have finished your antibiotic.

· Changes in sensation and possible nerve damage (Peripheral Neuropathy)

Damuge to the nerves in arms, hands, legs, or feet can happen in people taking fluoroquinolones, including levofloxacin. Talk with your healthcare provider right away if you get any of the following symptoms of peripheral neuropathy in your arms, hands, legs, or feet:

- o pain
- o burning
- o tingling
- o numbness

Levofloxacin tablets may need to be stopped to prevent permanent nerve damage.

• Serious heart rhythm changes (QT prolongation and torsades de pointes)

Tell your healthcare provider right away if you have a change in your heart beat (a fast or irregular heartbeat), or if you faint. Levofloxacin tablets may cause a rare heart problem known as prolongation of the QT interval. This condition can cause an abnormal heartbeat and can be very dangerous. The chances of this happening are higher in people:

- o who are elderly
- o with a family history of prolonged QT interval
- o with low blood potassium (hypokalemia)
- o who take certain medicines to control heart rhythm (antiarrhythmics)
- Joint Problems

Increased chance of problems with joints and tissues around joints in children can happen. Tell your child's healthcare provider if your child has any joint problems during or after treatment with level floxacin tables.

Changes in blood sugar

People who take levofloxacin tablets and other fluoroquimolone medicines with oral anti-diabetes medicines or with insulin can get low blood sugar (hypoglycemia) and high blood sugar (hypoglycemia) and high blood sugar (hyperglycemia). Follow your healthcare provider's instructions for how offen to check your blood sugar. If you have diabetes and you get low blood sugar while taking levofloxacin tablets, stop taking levofloxacin tablets and call your healthcare provider right away. Your antibiotic medicine may need to be changed.

• Sensitivity to sunlight (photosensitivity)

See "What should I avoid while taking levofloxacin tablets?"

The most common side effects of levofloxacin tablets include:

- diarrhea
- insomnia
- constipationdizziness

In children 6 months and older who take levofloxacin tablets to treat anthrax disease or plague, vomiting

Levofloxacin tablets may cause false-positive urine screening results for opiates when testing is done with some commercially available kits. A positive result should be confirmed using a more specific

These are not all the possible side effects of levofloxacin tablets. Tell your healthcare provider about any side effect that bothers you or that does not go away.

Call your doctor for medical advice about side effects. You may report side effects to FDA at 1-800-FDA-1088.

How should I store levofloxacin tablets?

Store at 20°-25°C (68°-77°F) [See USP Controlled Room Temperature]. Keep the container closed tightly.

Keep levofloxacin tablets and all medicines out of the reach of children.

General Information about the safe and effective use of levofloxacin tablets

Medicines are sometimes prescribed for purposes other than those listed in a Medication Guide. Do not use levofloxacin tablets for a condition for which it is not prescribed. Do not give levofloxacin tablets to other people, even if they have the same symptoms that you have. It may harm them.

This Medication Guide summarizes the most important information about levofloxacin tablets. If you would like more information about levofloxacin tablets, talk with your healthcare provider. You can ask your healthcare provider or pharmacist for information about levofloxacin tablets that is written for healthcare professionals.

For more information you can also call 1-800-346-6854.

What are the ingredients in levofloxacin tablets?

- 250 mg Levofloxacin Film-Coated Tablets:
- 250 lig Levolitoxatin min-Goateu i audies: Active ingredient: levofloxacin. Inactive ingredients: colloidal silicon dioxide, hypromellose, iron oxide red, magnesium stearate, microcrystalline cellulose, polyethylene glycol, polysorbate 80, povidone, propylene glycol, sodium starch glycolate and titanium dioxide.

- 500 mg Levofloxacin Film-Coated Tablets:
 Active ingredient: levofloxacin.
 Inactive ingredients: colloidal silicon dioxide, hypromellose, iron oxide red, iron oxide yellow, magnesium stearate, microcrystalline cellulose, polyethylene glycol, polysorbate 80, povidone, propylene glycol, sodium starch glycolate and titanium dioxide.
- 750 mg Levofloxacin Film-Coated Tablets:
- Active ingredient: levofloxacin.
 Inactive ingredients: colloidal silicon dioxide, hypromellose, magnesium stearate, microcrystalline cellulose, polyethylene glycol, polysorbate 80, povidone, propylene glycol, sodium starcl glycolate and titanium dioxide.

This Medication Guide has been approved by the U.S. Food and Drug Administration.

Manufactured by:

Wockhardt Limited,

Mumbai, India.

Distributed by:

Wockhardt USA LLC.

20 Waterview Blvd. Parsippany, NJ 07054

USA.

Rev.171012







LEVOFLOXACIN levofloxacin tablet					
Product Information					
Product Type	HUMAN PRESCRIPTION DRUG	Item Code (Source)	NDC:556	548-544
Route of Administration	ORAL				
Active Ingredient/Active	Moiety				
	Ingredient Name		Basis of	Strength	Strength
LEVOFLO XACIN (UNII: 6GNT3)	Ingredient Name (SLMF) (LEVOFLOXACIN ANHYDROUS - 1	JNII:RIX4E89 Y14)	LEVOFLO		250 mg
LEVOFLO XACIN (UNII: 6GNT3) Inactive Ingredients		JNIERIX4E89 Y14)			
		JNERIX4E89Y14)		DXACIN	
	/SLMF) (LEVOFLOXACIN ANHYDROUS - 1 Ingredient Name	JNII:RIX4E89 Y14)		DXACIN	250 mg
Inactive Ingredients	Ingredient Name LINE (UNIF OPER32261U)	JNIERIX4E89 Y14)		DXACIN	250 mg
Inactive Ingredients CELLULOSE, MICROCRYSTAL	.5LMF) (LEVOFLOXACIN ANHYDROUS - I Ingredient Name LLINE (UNIE OPIR32D61U) F3G675)	JNIERIX4E89Y14)		DXACIN	250 mg
Inactive Ingredients CELLULOSE, MICROCRYSTAL FERRIC O XIDE RED (UNIE 1K09	J.SLMF) (LEVOFLOXACIN ANNYDROUS - I Ingredient Name LLINE (UNIC OPIR32D6 IU) F36675) 29V3WO)	JNIERIX4E89Y14)		DXACIN	250 mg
Inactive Ingredients CELLULOSE, MICROCRYSTAI FERRIC OXIDE RED (UNII: 1K09 HYPROMELLOSES (UNII: 3NXW	JEME) (LEVOFLOXACIN ANHYDROUS - I Ingredient Name LLINE (UNIC OPIR32D6 IU) PEG675) 270977M6 IB0)	JNIERIX4E89Y14)		DXACIN	250 mg
Inactive Ingredients CELLULOSE, MICROCRYSTAL FERRIC OXIDE RED (UNIE 1K09 HYPROMELLOSES (UNIE 3KM) MAGNESIUM STEARATE (UNIE	JEMB) (LEVOFLOXACIN ANINYDROUS - I Ingredient Name LINE (UNIS OPIR32D6 IU) F260 575) 70097MB IBO) 393 ZEBIB)	JNIERIX4E89Y14)		DXACIN	250 mg
Inactive Ingredients CELLULOSE, MICRO CRYSTAI FERRIC OXIDE RED (UNIL 1K09 HYPRO MELLOSES (UNIL 1K09 MAGNESIUM STRAATE (UNIL POLYSORBATE 80 (UNIL 6OZE	JEMP) (LEVOFLOXACIN ANHYDROUS - I Ingredient Name LLINE (UNIC OPIR32D61U) P3G675) P3G9750 P3097M6 E0) 29 ZG81b)	UNIERIX4E89Y14)		DXACIN	250 mg
Inactive Ingredients CELLULOSE, MICROCRYSTAI FERRIC OXIDE RED (UNIE 1K09 HYPROMELLOSES (UNIE 3NXW MAGNESIUM STEARATE (UNIE POLYSORBATE 20 (UNIE 502 PO VIDONE (UNIE FZ989G184E)	Ingredient Name LINE (UNIS OF 1872 DE 1872	JNIERIX4E89Y14)		DXACIN	250 mg

DOLVETHYLENE GI	YCOL 400 (UNII: B697894SGQ)			
TOLILITIES OF	1002 400 (CML 203703430Q)			
Product Characte	eristics			
Color	PINK (Pink)		Score	no score
Shape	CAPSULE (Capsule-shaped)		Size	14mm
Flavor			Imprint Code	W;544
Contains				
Packaging				
# Item Code	Package Description	Marketii	ng Start Date	Marketing End Da
1 NDC:55648-544-01	20 in 1 BOTTLE			
2 NDC:55648-544-02	50 in 1 BOTTLE			
3 NDC:55648-544-03	100 in 1 BOTTLE			
4 NDC:55648-544-04	500 in 1 BOTTLE			
5 NDC:55648-544-05	100 in 1 CARTON			
5	10 in 1 BLISTER PACK			
Marketing Inf	ormation			
Marketing Categor		wanh Citation	Marketing Start I	Date Marketing End
	, Adminer of Monos	, up a Citation	markeding Start i	June Retting Ent
ANDA	ANDA090367		06/20/2011	

Marketing Catego	ory Applicati	on Number or Monogr	aph Citation	Marketing Sta	art Date	Marketing	End Date
ANDA	ANDA09036	7		06/20/2011			
EVOFLOXA	ACIN						
vofloxacin tablet							
Product Inforn	ation						
Product Type		HUMAN PRESCRIPTION	DRUG	Item Code (So	urce)	NDC:556	348-545
Route of Administ	ration	ORAL					
TOUR OF FRANKIS							
Active Ingredie	nt/Active Mo	ietv					
reure ingreun		Ingredient Name			Pacie of	Strength	Strongt
EVOEL O VACINO		F) (LEVOFLOXACIN ANI	IVDPOLIS - LINII-	DIYAERQ VIA)	LEVOFLO		500 mg
LEVOILOARCEN	JIME GOITTS ISEM	r) (EE TOTEO/EICHTHA	TIDIOUS - CIVIL	1004105114)	LLTOIL	JANGET .	Joo mg
nactive Ingred	ionto						
mactive mgreu	ients	Ingredient Na	ma				trength
TELLIII OSE MICI	OCDVSTALL IN	E (UNII: OP1R32D61U)	me			3	trengtn
ERRIC O XIDE REI							
FERRIC O XIDE YE							
HYPROMELLOSES							
MAGNESIUM STEA							
OLYSORBATE 80	(UNII: 6 OZP39 Z	58 H)					
PO VIDO NE (UNII: F	Z989GH94E)						
PRO PYLENE GLYO	OL (UNII: 6DC90	(167V3)					
SILICON DIO XIDE	(UNII: ETJ7Z6 XB)	J4)					
FITANIUM DIO XID	E (UNII: 15FIX9V2	JP)					
SODIUM STARCH	GLYCOLATE TY	PE A POTATO (UNII: 58	6J3G2A2)				
POLYETHYLENE C	LYCOL 400 (UN	II: B697894SGQ)					
Product Chara	teristics						
Color	ORANGE (Peach) Score		no sco	re			
Shape	CAPSULE (Cap	sule-shaped)	Size		18 mm		
Flavor				Imprint Code		W;545	
Contains							
Packaging							
# Item Cod	e Pac	kage Description	Marketin	g Start Date	Ma	rketing Er	d Date
1 NDC:55648-545-0	1 20 in 1 E	OTTLE					

Ħ	Item Code	Package Description	Marketti	ig Start Date	Marketing End Date
1	NDC:55648-545-01	20 in 1 BOTTLE			
2	NDC:55648-545-02	50 in 1 BOTTLE			
3	NDC:55648-545-03	100 in 1 BOTTLE			
4	NDC:55648-545-04	500 in 1 BOTTLE			
5	NDC:55648-545-05	100 in 1 CARTON			
5		10 in 1 BLISTER PACK			
N	Aarketing Info	rmation			
D	Marketing Category	Application Number or Monogra	aph Citation	Marketing Start Dat	te Marketing End Date

Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
ANDA	ANDA090367	06/20/2011	

LEVOFLOXACIN					
levo floxacin tablet					
Product Information					
Product Type	HUMAN PRESCRIPTION DRUG	Item Code (S	ource)	NDC:556	48-547
Route of Administration	ORAL				
Active Ingredient/Active M	Ioiety				
	Ingredient Name		Basis of S	trength	Streng
LEVOFLOXACIN (UNII: 6GNT3Y5	LMF) (LEVOFLOXACIN ANHYDROUS - U	NII:RIX4E89Y14)	LEVOFLOX	ACIN	750 mg
Inactive Ingredients					
	Ingredient Name			S	trength
CELLUL OCE MICROCRIVETALL	DIE CINIII ODIDADECHO				

mactive ingredients	
Ingredient Name	Strength
CELLULO SE, MICRO CRYSTALLINE (UNII: OPIR32D61U)	
HYPROMELLOSES (UNII: 3NXW29V3WO)	
MAGNESIUM STEARATE (UNII: 70 09 7M6 I30)	
POLYSORBATE 80 (UNII: 6 OZP39 ZG8 H)	
PO VIDO NE (UNII: FZ989GH94E)	
PROPYLENE GLYCOL (UNII: 6 DC9 Q167V3)	
SILICON DIO XIDE (UNII: ETJ7Z6 XBU4)	
TITANIUM DIO XIDE (UNII: 15FIX9 V2JP)	
SODIUM STARCH GLYCOLATE TYPE A POTATO (UNII: 5856 J3G2 A2)	
POLYETHYLENE GLYCOL 400 (UNI: B697894SGQ)	

Product Characteristics				
WHITE (White)	Score	no score		
CAPSULE (Capsule-shaped)	Size	22mm		
	Imprint Code	W;547		
	WHITE (White)	WHITE (White) Score CAPSULE (Capsule-shaped) Size		

P	ackaging			
#	Item Code	Package Description	Marketing Start Date	Marketing End Date
1	NDC:55648-547-01	20 in 1 BOTTLE		
2	NDC:55648-547-02	50 in 1 BOTTLE		
3	NDC:55648-547-03	100 in 1 BOTTLE		
4	NDC:55648-547-04	500 in 1 BOTTLE		
5	NDC:55648-547-05	100 in 1 CARTON		
5		10 in 1 BLISTER PACK		

Marketing Information				
Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date	
ANDA	ANDA090367	06/20/2011		

Labeler - Wockhardt Limited (650069115)

Registrant - Wockhardt Limited (650069115)

Estab	Establishment				
Name	Address	ID/FEI	Business Operations		
Wockhar	dt	676257570	ANALYSIS(55648-544, 55648-545, 55648-547), MANUFACTURE(55648-544, 55648-545, 55648-547), DACK(55648-544, 55648-545, 55648-547), LABEL(55648-544, 55648-545, 55648-547), LABEL(55648-545, 55648-545, 55648-547), LABEL(55648-544, 55648-545, 55648-547), LABEL(55648-545, 55648-547), LABEL(55648-545, 55648-547), LABEL(55648-545, 55648-545, 55648-547), LABEL(55648-545, 55648-545, 55648-547), LABEL(55648-545, 55648-55		

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