PRESCRIBING INFORMATION

2 **CEFTIN®** Tablets

3 (cefuroxime axetil tablets)

4

7

1

5 CEFTIN[®] for Oral Suspension

6 (cefuroxime axetil powder for oral suspension)

8 To reduce the development of drug-resistant bacteria and maintain the effectiveness of

9 CEFTIN and other antibacterial drugs, CEFTIN should be used only to treat or prevent infections

10 that are proven or strongly suspected to be caused by bacteria.

11 **DESCRIPTION**

12 CEFTIN Tablets and CEFTIN for Oral Suspension contain cefuroxime as cefuroxime axetil.

13 CEFTIN is a semisynthetic, broad-spectrum cephalosporin antibiotic for oral administration.

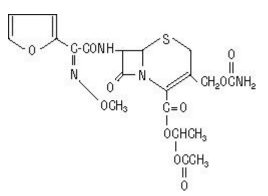
14 Chemically, cefuroxime axetil, the 1-(acetyloxy) ethyl ester of cefuroxime, is (*RS*)-1-

15 hydroxyethyl (6*R*,7*R*)-7-[2-(2-furyl)glyoxyl-amido]-3-(hydroxymethyl)-8-oxo-5-thia-1-

16 azabicyclo[4.2.0]-oct-2-ene-2-carboxylate, 7^2 -(Z)-(O-methyl-oxime), 1-acetate 3-carbamate. Its

- 17 molecular formula is $C_{20}H_{22}N_4O_{10}S$, and it has a molecular weight of 510.48.
- 18 Cefuroxime axetil is in the amorphous form and has the following structural formula:

19



20

21 CEFTIN Tablets are film-coated and contain the equivalent of 250 or 500 mg of cefuroxime

22 as cefuroxime axetil. CEFTIN Tablets contain the inactive ingredients colloidal silicon dioxide,

23 croscarmellose sodium, hydrogenated vegetable oil, hypromellose, methylparaben,

24 microcrystalline cellulose, propylene glycol, propylparaben, sodium benzoate, sodium lauryl

25 sulfate, and titanium dioxide.

26 CEFTIN for Oral Suspension, when reconstituted with water, provides the equivalent of

27 125 mg or 250 mg of cefuroxime (as cefuroxime axetil) per 5 mL of suspension. CEFTIN for

28 Oral Suspension contains the inactive ingredients acesulfame potassium, aspartame, povidone

29 K30, stearic acid, sucrose, tutti-frutti flavoring, and xanthan gum.

30

31 CLINICAL PHARMACOLOGY

- 32 **Absorption and Metabolism:** After oral administration, cefuroxime axetil is absorbed from
- 33 the gastrointestinal tract and rapidly hydrolyzed by nonspecific esterases in the intestinal mucosa
- 34 and blood to cefuroxime. Cefuroxime is subsequently distributed throughout the extracellular
- 35 fluids. The axetil moiety is metabolized to acetaldehyde and acetic acid.
- 36 Pharmacokinetics: Approximately 50% of serum cefuroxime is bound to protein. Serum
- 37 pharmacokinetic parameters for CEFTIN Tablets and CEFTIN for Oral Suspension are shown in
- 38 Tables 1 and 2.
- 39

40 **Table 1. Postprandial Pharmacokinetics of Cefuroxime Administered as CEFTIN Tablets**

41 to Adults*

Dose [†]	Peak Plasma	Time of Peak	Mean	
(Cefuroxime	Concentration	Plasma	Elimination	AUC
Equivalent)	(mcg/mL)	Concentration (hr)	Half-Life (hr)	(mcg-hr mL)
125 mg	2.1	2.2	1.2	6.7
250 mg	4.1	2.5	1.2	12.9
500 mg	7.0	3.0	1.2	27.4
1,000 mg	13.6	2.5	1.3	50.0

- 42 *Mean values of 12 healthy adult volunteers.
- 43 [†]Drug administered immediately after a meal.
- 44

45 **Table 2. Postprandial Pharmacokinetics of Cefuroxime Administered as CEFTIN for Oral**

46 Suspension to Pediatric Patients*

			Time of Peak	Mean	
$Dose^\dagger$		Peak Plasma	Plasma	Elimination	
(Cefuroxime		Concentration	Concentration	Half-Life	AUC
Equivalent)	n	(mcg/mL)	(hr)	(hr)	(mcg-hr mL)
10 mg/kg	8	3.3	3.6	1.4	12.4
15 mg/kg	12	5.1	2.7	1.9	22.5
20 mg/kg	8	7.0	3.1	1.9	32.8

47 *Mean age = 23 months.

- 48 [†]Drug administered with milk or milk products.
- 49
- 50 Comparative Pharmacokinetic Properties: A 250 mg/5 mL-dose of CEFTIN Suspension
- 51 is bioequivalent to 2 times 125 mg/5 mL-dose of CEFTIN Suspension when administered with
- 52 food (see Table 3). **CEFTIN for Oral Suspension was not bioequivalent to CEFTIN Tablets**
- 53 when tested in healthy adults. The tablet and powder for oral suspension formulations are
- 54 **NOT substitutable on a milligram-per-milligram basis.** The area under the curve for the
- 55 suspension averaged 91% of that for the tablet, and the peak plasma concentration for the
- 56 suspension averaged 71% of the peak plasma concentration of the tablets. Therefore, the safety

- 57 and effectiveness of both the tablet and oral suspension formulations had to be established in
- 58 separate clinical trials.
- 59

Table 3. Pharmacokinetics of Cefuroxime Administered as 250 mg/5 mL or 2 x 125 mg/5 mL CEFTIN for Oral Suspension to Adults* With Food

Dose	Peak Plasma	Time of Peak	Mean	
(Cefuroxime	Concentration	Plasma	Elimination	С
Equivalent)	(mcg/mL)	Concentration (hr)	Half-Life (hr)	(mcg-hr mL)
250 mg/5 mL	2.23	3	1.40	8.92
2 x 125 mg/5 mL	2.37	3	1.44	9.75

62 63

*Mean values of 18 healthy adult volunteers.

Food Effect on Pharmacokinetics: Absorption of the tablet is greater when taken after food
 (absolute bioavailability of CEFTIN Tablets increases from 37% to 52%). Despite this difference

66 in absorption, the clinical and bacteriologic responses of patients were independent of food

67 intake at the time of tablet administration in 2 studies where this was assessed.

All pharmacokinetic and clinical effectiveness and safety studies in pediatric patients using
 the suspension formulation were conducted in the fed state. No data are available on the

absorption kinetics of the suspension formulation when administered to fasted pediatric patients.

71 **Renal Excretion:** Cefuroxime is excreted unchanged in the urine; in adults, approximately

72 50% of the administered dose is recovered in the urine within 12 hours. The pharmacokinetics of

73 cefuroxime in the urine of pediatric patients have not been studied at this time. Until further data

are available, the renal pharmacokinetic properties of cefuroxime axetil established in adults

should not be extrapolated to pediatric patients.

76 Because cefuroxime is renally excreted, the serum half-life is prolonged in patients with

reduced renal function. In a study of 20 elderly patients (mean age = 83.9 years) having a mean

78 creatinine clearance of 34.9 mL/min, the mean serum elimination half-life was 3.5 hours. Despite

the lower elimination of cefuroxime in geriatric patients, dosage adjustment based on age is not

80 necessary (see PRECAUTIONS: Geriatric Use).

81 **Microbiology:** The in vivo bactericidal activity of cefuroxime axetil is due to cefuroxime's

82 binding to essential target proteins and the resultant inhibition of cell-wall synthesis.

83 Cefuroxime has bactericidal activity against a wide range of common pathogens, including

84 many beta-lactamase-producing strains. Cefuroxime is stable to many bacterial beta-lactamases,

85 especially plasmid-mediated enzymes that are commonly found in enterobacteriaceae.

86 Cefuroxime has been demonstrated to be active against most strains of the following

87 microorganisms both in vitro and in clinical infections as described in the INDICATIONS AND

88 USAGE section (see INDICATIONS AND USAGE section).

89 Aerobic Gram-Positive Microorganisms:

90 Staphylococcus aureus (including beta-lactamase-producing strains)

91 Streptococcus pneumoniae

92 Streptococcus pyogenes

93 Aerobic Gram-Negative Microorganisms:

- 94 Escherichia coli
- 95 *Haemophilus influenzae* (including beta-lactamase–producing strains)
- 96 Haemophilus parainfluenzae
- 97 Klebsiella pneumoniae
- 98 Moraxella catarrhalis (including beta-lactamase–producing strains)
- 99 *Neisseria gonorrhoeae* (including beta-lactamase–producing strains)

100 Spirochetes:

- 101 Borrelia burgdorferi
- 102 Cefuroxime has been shown to be active in vitro against most strains of the following
- 103 microorganisms; however, the clinical significance of these findings is unknown.
- 104 Cefuroxime exhibits in vitro minimum inhibitory concentrations (MICs) of 4.0 mcg/mL or
- 105 less (systemic susceptible breakpoint) against most (≥90%) strains of the following
- 106 microorganisms; however, the safety and effectiveness of cefuroxime in treating clinical
- 107 infections due to these microorganisms have not been established in adequate and
- 108 well-controlled trials.

109 Aerobic Gram-Positive Microorganisms:

- 110 Staphylococcus epidermidis
- 111 Staphylococcus saprophyticus
- 112 Streptococcus agalactiae
- 113 NOTE: *Listeria monocytogenes* and certain strains of enterococci, e.g., *Enterococcus faecalis*
- 114 (formerly *Streptococcus faecalis*), are resistant to cefuroxime. Methicillin-resistant staphylococci
- 115 are resistant to cefuroxime.

116 Aerobic Gram-Negative Microorganisms:

- 117 Morganella morganii
- 118 Proteus inconstans
- 119 Proteus mirabilis
- 120 Providencia rettgeri
- 121 NOTE: Pseudomonas spp., Campylobacter spp., Acinetobacter calcoaceticus, Legionella spp.,
- 122 and most strains of Serratia spp. and Proteus vulgaris are resistant to most first- and
- 123 second-generation cephalosporins. Some strains of Morganella morganii, Enterobacter cloacae,
- 124 and *Citrobacter* spp. have been shown by in vitro tests to be resistant to cefuroxime and other
- 125 cephalosporins.

126 Anaerobic Microorganisms:

- 127 Peptococcus niger
- 128 NOTE: Most strains of *Clostridium difficile* and *Bacteroides fragilis* are resistant to cefuroxime.
- 129 Susceptibility Tests: *Dilution Techniques:* Quantitative methods that are used to
- 130 determine MICs provide reproducible estimates of the susceptibility of bacteria to antimicrobial
- 131 compounds. One such standardized procedure uses a standardized dilution method¹ (broth, agar,

132 or microdilution) or equivalent with cefuroxime powder. The MIC values obtained should be

133 interpreted according to the following criteria:

134

MIC (mcg/mL)	Interpretation
≤4	(S) Susceptible
8-16	(I) Intermediate
≥32	(R) Resistant

135

136 A report of "Susceptible" indicates that the pathogen, if in the blood, is likely to be inhibited 137 by usually achievable concentrations of the antimicrobial compound in blood. A report of 138 "Intermediate" indicates that inhibitory concentrations of the antibiotic may be achieved if high 139 dosage is used or if the infection is confined to tissues or fluids in which high antibiotic 140 concentrations are attained. This category also provides a buffer zone that prevents small, 141 uncontrolled technical factors from causing major discrepancies in interpretation. A report of 142 "Resistant" indicates that usually achievable concentrations of the antimicrobial compound in the 143 blood are unlikely to be inhibitory and that other therapy should be selected. 144 Standardized susceptibility test procedures require the use of laboratory control 145 microorganisms. Standard cefuroxime powder should give the following MIC values:

146

<u>Microorganism</u>	MIC (mcg/mL)
Escherichia coli ATCC 25922	2-8
Staphylococcus aureus ATCC 29213	0.5-2

147

Diffusion Techniques: Quantitative methods that require measurement of zone diameters
 provide estimates of the susceptibility of bacteria to antimicrobial compounds. One such
 standardized procedure² that has been recommended (for use with disks) to test the susceptibility
 of microorganisms to cefuroxime uses the 30-mcg cefuroxime disk. Interpretation involves
 correlation of the diameter obtained in the disk test with the MIC for cefuroxime.
 Reports from the laboratory providing results of the standard single-disk susceptibility test
 with a 30-mcg cefuroxime disk should be interpreted according to the following criteria:

Interpretation
(S) Susceptible
(I) Intermediate
(R) Resistant

156

157 Interpretation should be as stated above for results using dilution techniques.

158 As with standard dilution techniques, diffusion methods require the use of laboratory control

159 microorganisms. The 30-mcg cefuroxime disk provides the following zone diameters in these

160 laboratory test quality control strains:

<u>Microorganism</u>	Zone Diameter (mm)
Escherichia coli ATCC 25922	20-26
Staphylococcus aureus ATCC 25923	27-35

162 INDICATIONS AND USAGE

163 NOTE: CEFTIN TABLETS AND CEFTIN FOR ORAL SUSPENSION ARE NOT

164 **BIOEQUIVALENT AND ARE NOT SUBSTITUTABLE ON A**

165 MILLIGRAM-PER-MILLIGRAM BASIS (SEE CLINICAL PHARMACOLOGY).

166 **CEFTIN Tablets:** CEFTIN Tablets are indicated for the treatment of patients with mild to

- 167 moderate infections caused by susceptible strains of the designated microorganisms in the
- 168 conditions listed below:
- 169 **1. Pharyngitis/Tonsillitis** caused by *Streptococcus pyogenes*.
- 170 **NOTE:** The usual drug of choice in the treatment and prevention of streptococcal infections,
- including the prophylaxis of rheumatic fever, is penicillin given by the intramuscular route.
- 172 CEFTIN Tablets are generally effective in the eradication of streptococci from the
- 173 nasopharynx; however, substantial data establishing the efficacy of cefuroxime in the
- subsequent prevention of rheumatic fever are not available. Please also note that in all clinical
- trials, all isolates had to be sensitive to both penicillin and cefuroxime. There are no data from
- adequate and well-controlled trials to demonstrate the effectiveness of cefuroxime in the
 treatment of penicillin-resistant strains of *Streptococcus pyogenes*.
- Acute Bacterial Otitis Media caused by *Streptococcus pneumoniae*, *Haemophilus influenzae* (including beta-lactamase–producing strains), *Moraxella catarrhalis* (including
 beta-lactamase–producing strains), or *Streptococcus pyogenes*.
- Acute Bacterial Maxillary Sinusitis caused by *Streptococcus pneumoniae* or *Haemophilus influenzae* (non-beta-lactamase–producing strains only). (See CLINICAL STUDIES section.)
- 183 **NOTE:** In view of the insufficient numbers of isolates of beta-lactamase–producing strains of
- 184 *Haemophilus influenzae* and *Moraxella catarrhalis* that were obtained from clinical trials with
- 185 CEFTIN Tablets for patients with acute bacterial maxillary sinusitis, it was not possible to
- adequately evaluate the effectiveness of CEFTIN Tablets for sinus infections known,
- suspected, or considered potentially to be caused by beta-lactamase–producing *Haemophilus*
- 188 *influenzae* or *Moraxella catarrhalis*.

189 4. Acute Bacterial Exacerbations of Chronic Bronchitis and Secondary Bacterial

- 190 Infections of Acute Bronchitis caused by *Streptococcus pneumoniae*, *Haemophilus*
- 191 *influenzae* (beta-lactamase negative strains), or *Haemophilus parainfluenzae* (beta-lactamase
- 192 negative strains). (See DOSAGE AND ADMINISTRATION section and CLINICAL
- 193STUDIES section.)
- 194 **5.** Uncomplicated Skin and Skin-Structure Infections caused by *Staphylococcus aureus* 195 (including beta-lactamase, producing strains) or *Strantococcus pyogenes*
- 195 (including beta-lactamase–producing strains) or *Streptococcus pyogenes*.

- 196 6. Uncomplicated Urinary Tract Infections caused by *Escherichia coli* or *Klebsiella* 197 pneumoniae. 198 7. Uncomplicated Gonorrhea, urethral and endocervical, caused by penicillinase-producing 199 and non-penicillinase-producing strains of Neisseria gonorrhoeae and uncomplicated 200 gonorrhea, rectal, in females, caused by non-penicillinase-producing strains of Neisseria 201 gonorrhoeae. 202 8. Early Lyme Disease (erythema migrans) caused by Borrelia burgdorferi. 203 204 **CEFTIN for Oral Suspension:** CEFTIN for Oral Suspension is indicated for the treatment of 205 pediatric patients 3 months to 12 years of age with mild to moderate infections caused by 206 susceptible strains of the designated microorganisms in the conditions listed below. The safety 207 and effectiveness of CEFTIN for Oral Suspension in the treatment of infections other than those 208 specifically listed below have not been established either by adequate and well-controlled trials 209 or by pharmacokinetic data with which to determine an effective and safe dosing regimen. 210 1. Pharyngitis/Tonsillitis caused by *Streptococcus pyogenes*. 211 **NOTE:** The usual drug of choice in the treatment and prevention of streptococcal infections, 212 including the prophylaxis of rheumatic fever, is penicillin given by the intramuscular route. 213 CEFTIN for Oral Suspension is generally effective in the eradication of streptococci from the 214 nasopharynx; however, substantial data establishing the efficacy of cefuroxime in the 215 subsequent prevention of rheumatic fever are not available. Please also note that in all clinical 216 trials, all isolates had to be sensitive to both penicillin and cefuroxime. There are no data from 217 adequate and well-controlled trials to demonstrate the effectiveness of cefuroxime in the 218 treatment of penicillin-resistant strains of Streptococcus pyogenes. 219 2. Acute Bacterial Otitis Media caused by Streptococcus pneumoniae, Haemophilus 220 influenzae (including beta-lactamase-producing strains), Moraxella catarrhalis (including 221 beta-lactamase-producing strains), or Streptococcus pyogenes. 222 3. Impetigo caused by Staphylococcus aureus (including beta-lactamase-producing strains) or 223 Streptococcus pyogenes. 224 225 To reduce the development of drug-resistant bacteria and maintain the effectiveness of 226 CEFTIN and other antibacterial drugs, CEFTIN should be used only to treat or prevent 227 infections that are proven or strongly suspected to be caused by susceptible bacteria. When 228 culture and susceptibility information are available, they should be considered in selecting or
 - 229 modifying antibacterial therapy. In the absence of such data, local epidemiology and
 - 230 susceptibility patterns may contribute to the empiric selection of therapy.

231 CONTRAINDICATIONS

CEFTIN products are contraindicated in patients with known allergy to the cephalosporingroup of antibiotics.

234 WARNINGS

235 **CEFTIN TABLETS AND CEFTIN FOR ORAL SUSPENSION ARE NOT** 236 **BIOEQUIVALENT AND ARE THEREFORE NOT SUBSTITUTABLE ON A** 237 MILLIGRAM-PER-MILLIGRAM BASIS (SEE CLINICAL PHARMACOLOGY). 238 **BEFORE THERAPY WITH CEFTIN PRODUCTS IS INSTITUTED, CAREFUL** 239 **INOUIRY SHOULD BE MADE TO DETERMINE WHETHER THE PATIENT HAS** 240 HAD PREVIOUS HYPERSENSITIVITY REACTIONS TO CEFTIN PRODUCTS, 241 **OTHER CEPHALOSPORINS, PENICILLINS, OR OTHER DRUGS. IF THIS** 242 **PRODUCT IS TO BE GIVEN TO PENICILLIN-SENSITIVE PATIENTS, CAUTION** 243 SHOULD BE EXERCISED BECAUSE CROSS-HYPERSENSITIVITY AMONG 244 BETA-LACTAM ANTIBIOTICS HAS BEEN CLEARLY DOCUMENTED AND MAY 245 **OCCUR IN UP TO 10% OF PATIENTS WITH A HISTORY OF PENICILLIN** 246 ALLERGY. IF A CLINICALLY SIGNIFICANT ALLERGIC REACTION TO CEFTIN 247 PRODUCTS OCCURS, DISCONTINUE THE DRUG AND INSTITUTE APPROPRIATE 248 THERAPY. SERIOUS ACUTE HYPERSENSITIVITY REACTIONS MAY REQUIRE 249 TREATMENT WITH EPINEPHRINE AND OTHER EMERGENCY MEASURES, 250 **INCLUDING OXYGEN, INTRAVENOUS FLUIDS, INTRAVENOUS** 251 ANTIHISTAMINES, CORTICOSTEROIDS, PRESSOR AMINES, AND AIRWAY 252 MANAGEMENT, AS CLINICALLY INDICATED. 253 *Clostridium difficile* associated diarrhea (CDAD) has been reported with use of nearly all

- antibacterial agents, including CEFTIN, 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.
- 263 If CDAD is suspected or confirmed, ongoing antibiotic use not directed against *C. difficile*264 may need to be discontinued. Appropriate fluid and electrolyte management, protein
 265
- supplementation, antibiotic treatment of *C. difficile*, and surgical evaluation should be instituted
- as clinically indicated.

267 **PRECAUTIONS**

- 268 **General:** As with other broad-spectrum antibiotics, prolonged administration of cefuroxime
- axetil may result in overgrowth of nonsusceptible microorganisms. If superinfection occurs
- 270 during therapy, appropriate measures should be taken.

- 271 Cephalosporins, including cefuroxime axetil, should be given with caution to patients
- 272 receiving concurrent treatment with potent diuretics because these diuretics are suspected of273 adversely affecting renal function.
- Cefuroxime axetil, as with other broad-spectrum antibiotics, should be prescribed with caution in individuals with a history of colitis. The safety and effectiveness of cefuroxime axetil have not been established in patients with gastrointestinal malabsorption. Patients with gastrointestinal malabsorption were excluded from participating in clinical trials of cefuroxime axetil.
- 279 Cephalosporins may be associated with a fall in prothrombin activity. Those at risk include 280 patients with renal or hepatic impairment or poor nutritional state, as well as patients receiving a 281 protracted course of antimicrobial therapy, and patients previously stabilized on anticoagulant 282 therapy. Prothrombin time should be monitored in patients at risk and exogenous Vitamin K 283 administered as indicated.
- Prescribing CEFTIN 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.
- 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 2 or more months after having taken the last dose of the antibiotic. If this occurs, patients should contact their physician as soon as possible.
- Information for Patients/Caregivers (Pediatric): *Phenylketonurics:* CEFTIN for Oral
 Suspension 125 mg/5 mL contains phenylalanine 11.8 mg per 5 mL (1 teaspoonful) constituted
 suspension. CEFTIN for Oral Suspension 250 mg/5 mL contains phenylalanine 25.2 mg per
 5 mL (1 teaspoonful) constituted suspension.
- During clinical trials, the tablet was tolerated by pediatric patients old enough to swallow the
 cefuroxime axetil tablet whole. The crushed tablet has a strong, persistent, bitter taste and
 should not be administered to pediatric patients in this manner. Pediatric patients who cannot
 swallow the tablet whole should receive the oral suspension.
- Discontinuation of therapy due to taste and/or problems of administering this drug occurred in 1.4% of pediatric patients given the oral suspension. Complaints about taste (which may impair compliance) occurred in 5% of pediatric patients.
- 303 3. Patients should be counseled that antibacterial drugs, including CEFTIN, should only be used
 304 to treat bacterial infections. They do not treat viral infections (e.g., the common cold). When
 305 CEFTIN is prescribed to treat a bacterial infection, patients should be told that although it is
- 306 common to feel better early in the course of therapy, the medication should be taken exactly
- 307 as directed. Skipping doses or not completing the full course of therapy may: (1) decrease the
- 308 effectiveness of the immediate treatment, and (2) increase the likelihood that bacteria will
- 309develop resistance and will not be treatable by CEFTIN or other antibacterial drugs in the
- 310 future.

- 311 **Drug/Laboratory Test Interactions:** A false-positive reaction for glucose in the urine may
- 312 occur with copper reduction tests (Benedict's or Fehling's solution or with CLINITEST[®] tablets),
- but not with enzyme-based tests for glycosuria (e.g., CLINISTIX[®]). As a false-negative result
- 314 may occur in the ferricyanide test, it is recommended that either the glucose oxidase or
- 315 hexokinase method be used to determine blood/plasma glucose levels in patients receiving
- cefuroxime axetil. The presence of cefuroxime does not interfere with the assay of serum and
- 317 urine creatinine by the alkaline picrate method.
- 318 **Drug/Drug Interactions:** Concomitant administration of probenecid with cefuroxime axetil
- tablets increases the area under the serum concentration versus time curve by 50%. The peak
- 320 serum cefuroxime concentration after a 1.5-g single dose is greater when taken with 1 g of
- 321 probenecid (mean = 14.8 mcg/mL) than without probenecid (mean = 12.2 mcg/mL).
- 322 Drugs that reduce gastric acidity may result in a lower bioavailability of CEFTIN compared 323 with that of fasting state and tend to cancel the effect of postprandial absorption.
- In common with other antibiotics, cefuroxime axetil may affect the gut flora, leading to lower
 estrogen reabsorption and reduced efficacy of combined oral estrogen/progesterone
- 326 contraceptives.
- 327 Carcinogenesis, Mutagenesis, Impairment of Fertility: Although lifetime studies in
- 328 animals have not been performed to evaluate carcinogenic potential, no mutagenic activity was
- 329 found for cefuroxime axetil in a battery of bacterial mutation tests. Positive results were obtained
- in an in vitro chromosome aberration assay; however, negative results were found in an in vivo
- 331 micronucleus test at doses up to 1.5 g/kg. Reproduction studies in rats at doses up to
- 1,000 mg/kg/day (9 times the recommended maximum human dose based on mg/m²) have
- 333 revealed no impairment of fertility.
- 334 Pregnancy: Teratogenic Effects: Pregnancy Category B. Reproduction studies have been 335 performed in mice at doses up to 3,200 mg/kg/day (14 times the recommended maximum human 336 dose based on mg/m²) and in rats at doses up to 1,000 mg/kg/day (9 times the recommended
- dose based on mg/m^2) and in rats at doses up to 1,000 mg/kg/day (9 times the recommended maximum human dose based on mg/m^2) and have revealed no evidence of impaired fertility or
- harm to the fetus due to cefuroxime axetil. There are, however, no adequate and well-controlled
- 339 studies in pregnant women. Because animal reproduction studies are not always predictive of
- 340 human response, this drug should be used during pregnancy only if clearly needed.
- 341 **Labor and Delivery:** Cefuroxime axetil has not been studied for use during labor and delivery.
- 342 **Nursing Mothers:** Because cefuroxime is excreted in human milk, consideration should be
- 343 given to discontinuing nursing temporarily during treatment with cefuroxime axetil.
- 344 **Pediatric Use:** The safety and effectiveness of CEFTIN have been established for pediatric
- 345 patients aged 3 months to 12 years for acute bacterial maxillary sinusitis based upon its approval
- 346 in adults. Use of CEFTIN in pediatric patients is supported by pharmacokinetic and safety data in
- 347 adults and pediatric patients, and by clinical and microbiological data from adequate and
- 348 well-controlled studies of the treatment of acute bacterial maxillary sinusitis in adults and of
- 349 acute otitis media with effusion in pediatric patients. It is also supported by postmarketing
- 350 adverse events surveillance (see CLINICAL PHARMACOLOGY, INDICATIONS AND

351 USAGE, ADVERSE REACTIONS, DOSAGE AND ADMINISTRATION, and CLINICAL352 STUDIES).

353 **Geriatric Use:** Of the total number of subjects who received cefuroxime axetil in 20 clinical 354 studies of CEFTIN, 375 were 65 and over while 151 were 75 and over. No overall differences in

355 safety or effectiveness were observed between these subjects and younger adult subjects. The

356 geriatric patients reported somewhat fewer gastrointestinal events and less frequent vaginal

357 candidiasis compared with patients aged 12 to 64 years old; however, no clinically significant

358 differences were reported between the elderly and younger adult patients. Other reported clinical

- 359 experience has not identified differences in responses between the elderly and younger adult
- 360 patients.

361 **ADVERSE REACTIONS**

362 CEFTIN TABLETS IN CLINICAL TRIALS: Multiple-Dose Dosing Regimens: 7 to

10 Days Dosing: Using multiple doses of cefuroxime axetil tablets, 912 patients were treated
with cefuroxime axetil (125 to 500 mg twice daily). There were no deaths or permanent

365 disabilities thought related to drug toxicity. Twenty (2.2%) patients discontinued medication due 366 to adverse events thought by the investigators to be possibly, probably, or almost certainly

related to drug toxicity. Seventeen (85%) of the 20 patients who discontinued therapy did so

368 because of gastrointestinal disturbances, including diarrhea, nausea, vomiting, and abdominal

369 pain. The percentage of cefuroxime axetil tablet-treated patients who discontinued study drug

because of adverse events was very similar at daily doses of 1,000, 500, and 250 mg (2.3%,

2.1%, and 2.2%, respectively). However, the incidence of gastrointestinal adverse events

increased with the higher recommended doses.

373 The following adverse events were thought by the investigators to be possibly, probably, or

almost certainly related to cefuroxime axetil tablets in multiple-dose clinical trials (n = 912

375 cefuroxime axetil-treated patients).

376

Incidence ≥1%	Diarrhea/loose stools	3.7%		
	Nausea/vomiting	3.0%		
	Transient elevation in AST	2.0%		
	Transient elevation in ALT	1.6%		
	Eosinophilia	1.1%		
	Transient elevation in LDH	1.0%		
Incidence	Abdominal pain			
<1% but >0.1%	Abdominal cramps			
	Flatulence	_		
	Indigestion			
	Headache			
	Vaginitis			
	Vulvar itch			
	Rash			
	Hives			
	Itch			
	Dysuria			
	Chills			
	Chest pain			
	Shortness of breath			
	Mouth ulcers			
	Swollen tongue			
	Sleepiness			
	Thirst			
	Anorexia			
	Positive Coombs test			

377 Table 4. Adverse Reactions—CEFTIN Tablets

378 Multiple-Dose Dosing Regimens—Clinical Trials

379

5-Day Experience (see CLINICAL STUDIES section): In clinical trials using CEFTIN
 in a dose of 250 mg twice daily in the treatment of secondary bacterial infections of acute
 bronchitis, 399 patients were treated for 5 days and 402 patients were treated for 10 days. No
 difference in the occurrence of adverse events was found between the 2 regimens.

In Clinical Trials for Early Lyme Disease With 20 Days Dosing: Two multicenter
 trials assessed cefuroxime axetil tablets 500 mg twice a day for 20 days. The most common
 drug-related adverse experiences were diarrhea (10.6% of patients), Jarisch-Herxheimer reaction
 (5.6%), and vaginitis (5.4%). Other adverse experiences occurred with frequencies comparable
 to those reported with 7 to 10 days dosing.

389 Single-Dose Regimen for Uncomplicated Gonorrhea: In clinical trials using a single

dose of cefuroxime axetil tablets, 1,061 patients were treated with the recommended dosage of

- 391 cefuroxime axetil (1,000 mg) for the treatment of uncomplicated gonorrhea. There were no
- 392 deaths or permanent disabilities thought related to drug toxicity in these studies.
- 393 The following adverse events were thought by the investigators to be possibly, probably, or
- 394 almost certainly related to cefuroxime axetil in 1,000-mg single-dose clinical trials of
- 395 cefuroxime axetil tablets in the treatment of uncomplicated gonorrhea conducted in the United
- 396 States.
- 397

398Table 5. Adverse Reactions—CEFTIN Tablets

399 1-g Single-Dose Regimen for Uncomplicated Gonorrhea—Clinical Trials

Incidence ≥1%	Nausea/vomiting 6.8%	
	Diarrhea 4.2%	
Incidence	Abdominal pain	
<1% but >0.1%	Dyspepsia	
	Erythema	
	Rash	
	Pruritus	
	Vaginal candidiasis	
	Vaginal itch	
	Vaginal discharge	
	Headache	
	Dizziness	
	Somnolence	
	Muscle cramps	
	Muscle stiffness	
	Muscle spasm of neck	
	Tightness/pain in chest	
	Bleeding/pain in urethra	
	Kidney pain	
	Tachycardia	
	Lockjaw-type reaction	

400 CEFTIN FOR ORAL SUSPENSION IN CLINICAL TRIALS

401 In clinical trials using multiple doses of cefuroxime axetil powder for oral suspension,

402 pediatric patients (96.7% of whom were younger than 12 years of age) were treated with the

403 recommended dosages of cefuroxime axetil (20 to 30 mg/kg/day divided twice a day up to a

404 maximum dose of 500 or 1,000 mg/day, respectively). There were no deaths or permanent

405 disabilities in any of the patients in these studies. Eleven US patients (1.2%) discontinued

406 medication due to adverse events thought by the investigators to be possibly, probably, or almost

407 certainly related to drug toxicity. The discontinuations were primarily for gastrointestinal

408 disturbances, usually diarrhea or vomiting. During clinical trials, discontinuation of therapy due

- 409 to the taste and/or problems with administering this drug occurred in 13 (1.4%) pediatric patients
- 410 enrolled at centers in the United States.
- 411 The following adverse events were thought by the investigators to be possibly, probably, or
- 412 almost certainly related to cefuroxime axetil for oral suspension in multiple-dose clinical trials
- 413 (n = 931 cefuroxime axetil-treated US patients).
- 414

415 **Table 6. Adverse Reactions—CEFTIN for Oral Suspension**

416 Multiple-Dose Dosing Regimens—Clinical Trials

The state of the s		
Incidence $\geq 1\%$	Diarrhea/loose stools 8.6%	
	Dislike of taste 5.0%	
	Diaper rash 3.4%	
	Nausea/vomiting 2.6%	
Incidence	Abdominal pain	
<1% but >0.1%	Flatulence	
	Gastrointestinal infection	
	Candidiasis	
	Vaginal irritation	
	Rash	
	Hyperactivity	
	Irritable behavior	
	Eosinophilia	
	Positive direct Coombs test	
	Elevated liver enzymes	
	Viral illness	
	Upper respiratory infection	
	Sinusitis	
	Cough	
	Urinary tract infection	
	Joint swelling	
	Arthralgia	
	Fever	
	Ptyalism	

417 **POSTMARKETING EXPERIENCE WITH CEFTIN PRODUCTS**

- 418 In addition to adverse events reported during clinical trials, the following events have been
- 419 identified during clinical practice in patients treated with CEFTIN Tablets or with CEFTIN for
- 420 Oral Suspension and were reported spontaneously. Data are generally insufficient to allow an
- 421 estimate of incidence or to establish causation.
- 422 *General:* The following hypersensitivity reactions have been reported: anaphylaxis,
- 423 angioedema, pruritus, rash, serum sickness-like reaction, urticaria.

- 424 *Gastrointestinal:* Pseudomembranous colitis (see WARNINGS).
- 425 *Hematologic:* Hemolytic anemia, leukopenia, pancytopenia, thrombocytopenia, and
- 426 increased prothrombin time.
- 427 *Hepatic:* Hepatic impairment including hepatitis and cholestasis, jaundice.
- 428 **Neurologic:** Seizure.
- 429 **Skin:** Erythema multiforme, Stevens-Johnson syndrome, toxic epidermal necrolysis.
- 430 *Urologic:* Renal dysfunction.

431 CEPHALOSPORIN-CLASS ADVERSE REACTIONS

- 432 In addition to the adverse reactions listed above that have been observed in patients treated
- 433 with cefuroxime axetil, the following adverse reactions and altered laboratory tests have been
- 434 reported for cephalosporin-class antibiotics: toxic nephropathy, aplastic anemia, hemorrhage,
- 435 increased BUN, increased creatinine, false-positive test for urinary glucose, increased alkaline
- 436 phosphatase, neutropenia, elevated bilirubin, and agranulocytosis.
- 437 Several cephalosporins have been implicated in triggering seizures, particularly in patients
- 438 with renal impairment when the dosage was not reduced (see DOSAGE AND
- 439 ADMINISTRATION and OVERDOSAGE). If seizures associated with drug therapy occur, the
- 440 drug should be discontinued. Anticonvulsant therapy can be given if clinically indicated.

441 **OVERDOSAGE**

- 442 Overdosage of cephalosporins can cause cerebral irritation leading to convulsions. Serum
- 443 levels of cefuroxime can be reduced by hemodialysis and peritoneal dialysis.

444 DOSAGE AND ADMINISTRATION

- 445 NOTE: CEFTIN TABLETS AND CEFTIN FOR ORAL SUSPENSION ARE NOT
- 446 BIOEQUIVALENT AND ARE NOT SUBSTITUTABLE ON A
- 447 MILLIGRAM-PER-MILLIGRAM BASIS (SEE CLINICAL PHARMACOLOGY).
- 448

449 **Table 7. CEFTIN Tablets**

450 (May be administered without regard to meals.)

Dosage	Duration (days)			
Dosuge				
250 mg b.i.d.	10			
250 mg b.i.d.	10			
250 or 500 mg b.i.d.	10*			
250 or 500 mg b.i.d.	5-10			
250 or 500 mg b.i.d.	10			
250 mg b.i.d.	7-10			
1,000 mg once	single dose			
500 mg b.i.d.	20			
Pediatric Patients (who can swallow tablets whole)				
250 mg b.i.d.	10			
250 mg b.i.d.	10			
	250 mg b.i.d. 250 or 500 mg b.i.d. 250 or 500 mg b.i.d. 250 or 500 mg b.i.d. 250 mg b.i.d. 1,000 mg once 500 mg b.i.d. 250 mg b.i.d.			

* The safety and effectiveness of CEFTIN administered for less than 10 days in patients with
 acute exacerbations of chronic bronchitis have not been established.

- 453
- 454 **CEFTIN for Oral Suspension:** CEFTIN for Oral Suspension may be administered to
- 455 pediatric patients ranging in age from 3 months to 12 years, according to dosages in Table 8:
- 456

457 Table 8. CEFTIN for Oral Suspension

458 (Must be administered with food. Shake well each time before using.)

		Daily		
		Maximum	Duration	
Population/Infection	Dosage	Dose	(days)	
Pediatric Patients (3 months to 12 years)				
Pharyngitis/tonsillitis	20 mg/kg/day divided b.i.d.	500 mg	10	
Acute otitis media	30 mg/kg/day divided b.i.d.	1,000 mg	10	
Acute bacterial maxillary sinusitis	30 mg/kg/day divided b.i.d.	1,000 mg	10	
Impetigo	30 mg/kg/day divided b.i.d.	1,000 mg	10	

459

460 **Patients With Renal Failure:** The safety and efficacy of cefuroxime axetil in patients with

461 renal failure have not been established. Since cefuroxime is renally eliminated, its half-life will

462 be prolonged in patients with renal failure.

464 dispensing as follows:

- 465 1. Shake the bottle to loosen the powder.
- 466 2. Remove the cap.

⁴⁶³ **Directions for Mixing CEFTIN for Oral Suspension:** Prepare a suspension at the time of

- 467 3. Add the total amount of water for reconstitution (see Table 9) and replace the cap.
- 4684. Invert the bottle and vigorously rock the bottle from side to side so that water rises through469the powder.
- 470 5. Once the sound of the powder against the bottle disappears, turn the bottle upright and
- 471 vigorously shake it in a diagonal direction.
- 472

473 Table 9. Amount of Water Required for Reconstitution of Labeled Volumes of CEFTIN for 474 Oral Suspension

CEFTIN for Oral Suspension	Labeled Volume After Reconstitution	Amount of Water Required for Reconstitution
125 mg/5 mL	100 mL	37 mL
250 mg/5 mL	50 mL	19 mL
230 mg/3 mL	100 mL	35 mL

475

476 NOTE: SHAKE THE ORAL SUSPENSION WELL BEFORE EACH USE. Replace cap

477 securely after each opening. Store the reconstituted suspension between 2° and 8°C (36° and

478 46°F) (in a refrigerator). DISCARD AFTER 10 DAYS.

479 HOW SUPPLIED

- 480 **CEFTIN Tablets:** CEFTIN Tablets, 250 mg of cefuroxime (as cefuroxime axetil), are white,
- 481 capsule-shaped, film-coated tablets engraved with "GX ES7" on one side and blank on the other482 side as follows:
- 483 20 Tablets/Bottle NDC 0173-0387-00
- 484 CEFTIN Tablets, 500 mg of cefuroxime (as cefuroxime axetil), are white, capsule-shaped,

485 film-coated tablets engraved with "GX EG2" on one side and blank on the other side as follows:

- 486 20 Tablets/Bottle NDC 0173-0394-00
- 487 60 Tablets/Bottle NDC 0173-0394-42
- 488 Store the tablets between 15° and 30°C (59° and 86°F). Replace cap securely after each 489 opening.
- 490 **CEFTIN for Oral Suspension:** CEFTIN for Oral Suspension is provided as dry, white to
- 491 off-white, tutti-frutti-flavored powder. When reconstituted as directed, CEFTIN for Oral
- 492 Suspension provides the equivalent of 125 mg or 250 mg of cefuroxime (as cefuroxime axetil)
- 493 per 5 mL of suspension. It is supplied in amber glass bottles as follows:
- 494 **125 mg/5 mL:**
- 495 100-mL Suspension NDC 0173-0740-00
- 496 **250 mg/5 mL**:
- 497 50-mL Suspension NDC 0173-0741-10
- 498 100-mL Suspension NDC 0173-0741-00
- 499 Before reconstitution, store dry powder between 2° and 30°C (36° and 86°F).

500 After reconstitution, immediately store suspension between 2° and 8°C (36° and 46°F), 501 in a refrigerator. DISCARD AFTER 10 DAYS.

502 CLINICAL STUDIES

503 Ceftin Tablets: Acute Bacterial Maxillary Sinusitis: One adequate and well-controlled

study was performed in patients with acute bacterial maxillary sinusitis. In this study each

505 patient had a maxillary sinus aspirate collected by sinus puncture before treatment was initiated 506 for presumptive acute bacterial sinusitis. All patients had to have radiographic and clinical

- 507 evidence of acute maxillary sinusitis. As shown in the following summary of the study, the
- 508 general clinical effectiveness of CEFTIN Tablets was comparable to an oral antimicrobial agent
- 509 that contained a specific beta-lactamase inhibitor in treating acute maxillary sinusitis. However,
- 510 sufficient microbiology data were obtained to demonstrate the effectiveness of CEFTIN Tablets
- 511 in treating acute bacterial maxillary sinusitis due only to Streptococcus pneumoniae or
- 512 non-beta-lactamase-producing Haemophilus influenzae. An insufficient number of
- 513 beta-lactamase-producing *Haemophilus influenzae* and *Moraxella catarrhalis* isolates were

514 obtained in this trial to adequately evaluate the effectiveness of CEFTIN Tablets in the treatment

- 515 of acute bacterial maxillary sinusitis due to these 2 organisms.
- 516 This study enrolled 317 adult patients, 132 patients in the United States and 185 patients in

517 South America. Patients were randomized in a 1:1 ratio to cefuroxime axetil 250 mg twice daily

518 or an oral antimicrobial agent that contained a specific beta-lactamase inhibitor. An

- 519 intent-to-treat analysis of the submitted clinical data yielded the following results:
- 520

521 **Table 10. Clinical Effectiveness of CEFTIN Tablets Compared to Beta-Lactamase**

522 Inhibitor-Containing Control Drug in the Treatment of Acute Bacterial Maxillary Sinusitis

	US Patients*		South American Patients ^{\dagger}	
	CEFTIN	Control	CEFTIN	Control
	(n = 49)	(n = 43)	(n = 87)	(n = 89)
Clinical success (cure + improvement)	65%	53%	77%	74%
Clinical cure	53%	44%	72%	64%
Clinical improvement	12%	9%	5%	10%

523 * 95% Confidence interval around the success difference [-0.08, +0.32].

524 ^{\dagger} 95% Confidence interval around the success difference [-0.10, +0.16].

525

526 In this trial and in a supporting maxillary puncture trial, 15 evaluable patients had

- 527 non-beta-lactamase-producing *Haemophilus influenzae* as the identified pathogen. Ten (10) of
- 528 these 15 patients (67%) had their pathogen (non-beta-lactamase-producing Haemophilus

529 *influenzae*) eradicated. Eighteen (18) evaluable patients had *Streptococcus pneumoniae* as the

530 identified pathogen. Fifteen (15) of these 18 patients (83%) had their pathogen (*Streptococcus*

531 *pneumoniae*) eradicated.

- 532 **Safety:** The incidence of drug-related gastrointestinal adverse events was statistically
- 533 significantly higher in the control arm (an oral antimicrobial agent that contained a specific
- beta-lactamase inhibitor) versus the cefuroxime axetil arm (12% versus 1%, respectively;
- 535 P<.001), particularly drug-related diarrhea (8% versus 1%, respectively; P = .001).
- *Early Lyme Disease:* Two adequate and well-controlled studies were performed in patients
 with early Lyme disease. In these studies all patients had to present with physician-documented
 erythema migrans, with or without systemic manifestations of infection. Patients were
 randomized in a 1:1 ratio to a 20-day course of treatment with cefuroxime axetil 500 mg twice
 daily or doxycycline 100 mg 3 times daily. Patients were assessed at 1 month posttreatment for
 success in treating early Lyme disease (Part I) and at 1 year posttreatment for success in
 preventing the progression to the sequelae of late Lyme disease (Part II).
- 543 A total of 355 adult patients (181 treated with cefuroxime axetil and 174 treated with
- 544 doxycycline) were enrolled in the 2 studies. In order to objectively validate the clinical diagnosis
- of early Lyme disease in these patients, 2 approaches were used: 1) blinded expert reading of
- 546 photographs, when available, of the pretreatment erythema migrans skin lesion; and 2) serologic
- 547 confirmation (using enzyme-linked immunosorbent assay [ELISA] and immunoblot assay
- 548 ["Western" blot]) of the presence of antibodies specific to *Borrelia burgdorferi*, the etiologic
- agent of Lyme disease. By these procedures, it was possible to confirm the physician diagnosis
- of early Lyme disease in 281 (79%) of the 355 study patients. The efficacy data summarized
- below are specific to this "validated" patient subset, while the safety data summarized below
- reflect the entire patient population for the 2 studies.
- Analysis of the submitted clinical data for evaluable patients in the "validated" patient subsetyielded the following results:
- 555

556 Table 11. Clinical Effectiveness of CEFTIN Tablets Compared to Doxycycline in the

557 Treatment of Early Lyme Disease

	Pa	art I	Part II		
	(1 Month Pc	sttreatment)*	$(1 \text{ Year Posttreatment})^{\dagger}$		
	CEFTIN	Doxycycline	CEFTIN	Doxycycline	
	(n = 125)	(n = 108)	$(n = 105^{\ddagger})$	$(n = 83^{\ddagger})$	
Satisfactory clinical outcome [§]	91%	93%	84%	87%	
Clinical cure/success	72%	73%	73%	73%	
Clinical improvement	19%	19%	10%	13%	

- * 95% confidence interval around the satisfactory difference for Part I (-0.08, +0.05).
- 559 [†] 95% confidence interval around the satisfactory difference for Part II (-0.13, +0.07).
- 560 [‡] n's include patients assessed as unsatisfactory clinical outcomes (failure + recurrence) in
- 561 Part I (CEFTIN 11 [5 failure, 6 recurrence]; doxycycline 8 [6 failure, 2 recurrence]).
- 562 [§] Satisfactory clinical outcome includes cure + improvement (Part I) and success +

563 improvement (Part II).

564

565 CEFTIN and doxycycline were effective in prevention of the development of sequelae of late566 Lyme disease.

567 Safety: Drug-related adverse events affecting the skin were reported significantly more 568 frequently by patients treated with doxycycline than by patients treated with cefuroxime axetil 569 (12% versus 3%, respectively; P = .002), primarily reflecting the statistically significantly higher 570 incidence of drug-related photosensitivity reactions in the doxycycline arm versus the 571 cefuroxime axetil arm (9% versus 0%, respectively; P<.001). While the incidence of drug-related 572 gastrointestinal adverse events was similar in the 2 treatment groups (cefuroxime axetil - 13%; 573 doxycycline - 11%), the incidence of drug-related diarrhea was statistically significantly higher 574 in the cefuroxime axetil arm versus the doxycycline arm (11% versus 3%, respectively; 575 P = .005).

576 Secondary Bacterial Infections of Acute Bronchitis: Four randomized, controlled 577 clinical studies were performed comparing 5 days versus 10 days of CEFTIN for the treatment of 578 patients with secondary bacterial infections of acute bronchitis. These studies enrolled a total of 579 1,253 patients (CAE-516 n = 360; CAE-517 n = 177; CAEA4001 n = 362; CAEA4002 n = 354). The protocols for CAE-516 and CAE-517 were identical and compared CEFTIN 250 mg twice 580 daily for 5 days, CEFTIN 250 mg twice daily for 10 days, and AUGMENTIN[®] 500 mg 3 times 581 daily for 10 days. These 2 studies were conducted simultaneously. CAEA4001 and CAEA4002 582 583 compared CEFTIN 250 mg twice daily for 5 days, CEFTIN 250 mg twice daily for 10 days, and CECLOR[®] 250 mg 3 times daily for 10 days. They were otherwise identical to CAE-516 and 584 585 CAE-517 and were conducted over the following 2 years. Patients were required to have 586 polymorphonuclear cells present on the Gram stain of their screening sputum specimen, but 587 isolation of a bacterial pathogen from the sputum culture was not required for inclusion. The 588 following table demonstrates the results of the clinical outcome analysis of the pooled studies 589 CAE-516/CAE-517 and CAEA4001/CAEA4002, respectively:

590

591 Table 12. Clinical Effectiveness of CEFTIN Tablets 250 mg Twice Daily in Secondary

- 592 Bacterial Infections of Acute Bronchitis: Comparison of 5 Versus 10 Days' Treatment
- 593 **Duration**

	CAE-516 and CAE-517*		CAEA4001 and CAEA4002 ^{\dagger}	
	5 Day 10 Day		5 Day	10 Day
	(n = 127)	(n = 139)	(n = 173)	(n = 192)
Clinical success (cure + improvement)	80%	87%	84%	82%
Clinical cure	61%	70%	73%	72%
Clinical improvement	19%	17%	11%	10%

* 95% Confidence interval around the success difference [-0.164, +0.029].

595 ^{\dagger} 95% Confidence interval around the success difference [-0.061, +0.103].

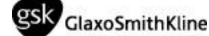
596

597 The response rates for patients who were both clinically and bacteriologically evaluable were 598 consistent with those reported for the clinically evaluable patients.

- 599 **Safety:** In these clinical trials, 399 patients were treated with CEFTIN for 5 days and
- 600 402 patients with CEFTIN for 10 days. No difference in the occurrence of adverse events was
- 601 observed between the 2 regimens.

602 **REFERENCES**

- National Committee for Clinical Laboratory Standards. *Methods for Dilution Antimicrobial Susceptibility Tests for Bacteria that Grow Aerobically*. 3rd ed. Approved Standard NCCLS
 Document M7-A3, Vol. 13, No. 25. Villanova, Pa: NCCLS; 1993.
- 806 2. National Committee for Clinical Laboratory Standards. *Performance Standards for*807 *Antimicrobial Disk Susceptibility Tests*. 4th ed. Approved Standard NCCLS Document
 808 M2-A4, Vol. 10, No. 7. Villanova, Pa: NCCLS; 1990.
- 609



- 610 611
- 612 GlaxoSmithKline
- 613 Research Triangle Park, NC 27709
- 614
- 615 CEFTIN is a registered trademark of GlaxoSmithKline.
- 616 CLINITEST and CLINISTIX are registered trademarks of Ames Division, Miles Laboratories,
- 617 Inc.
- 618
- 619 ©2007, GlaxoSmithKline
- 620 All rights reserved.
- 621
- 622 January 2007

RL-2353