

## PRESCRIBING INFORMATION

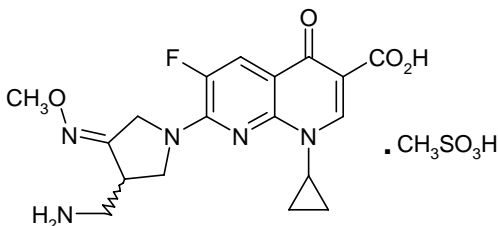
### FACTIVE<sup>®</sup> (gemifloxacin mesylate) Tablets

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

#### DESCRIPTION

FACTIVE (gemifloxacin mesylate) is a synthetic broad-spectrum antibacterial agent for oral administration. Gemifloxacin, a compound related to the fluoroquinolone class of antibiotics, is available as the mesylate salt in the sesquihydrate form. Chemically, gemifloxacin is (*R,S*)-7-[(4*Z*)-3-(aminomethyl)-4-(methoxyimino)-1-pyrrolidinyl]-1-cyclopropyl-6-fluoro-1,4-dihydro-4-oxo-1,8-naphthyridine-3-carboxylic acid.

The mesylate salt is a white to light brown solid with a molecular weight of 485.49. Gemifloxacin is considered freely soluble at neutral pH (350 µg/mL at 37°C, pH 7.0). Its empirical formula is C<sub>18</sub>H<sub>20</sub>FN<sub>5</sub>O<sub>4</sub>•CH<sub>4</sub>O<sub>3</sub>S and its chemical structure is:



Each white to off-white, oval, film-coated FACTIVE tablet has breaklines and GE 320 debossed on both faces and contains gemifloxacin mesylate equivalent to 320 mg gemifloxacin. The inactive ingredients are crospovidone, hydroxypropyl methylcellulose, magnesium stearate, microcrystalline cellulose, polyethylene glycol, povidone, and titanium dioxide.

#### CLINICAL PHARMACOLOGY

##### *Pharmacokinetics*

The pharmacokinetics of gemifloxacin are approximately linear over the dose range from 40 mg to 640 mg. There was minimal accumulation of gemifloxacin following multiple oral doses up to 640 mg a day for 7 days (mean accumulation <20%). Following repeat oral administration of 320 mg gemifloxacin once daily, steady-state is achieved by the third day of dosing.

##### *Absorption and Bioavailability*

Gemifloxacin, given as an oral tablet, is rapidly absorbed from the gastrointestinal tract. Peak plasma concentrations of gemifloxacin were observed between 0.5 and 2 hours following oral tablet administration and the absolute bioavailability of the 320 mg tablet averaged approximately 71% (95% CI 60%-84%). Following repeat oral doses of 320 mg to healthy subjects, the mean ± SD maximal gemifloxacin plasma concentrations

42 (C<sub>max</sub>) and systemic drug exposure (AUC (0-24)) were 1.61 ± 0.51 µg/mL (range 0.70-  
43 2.62 µg/mL) and 9.93 ± 3.07 µg•hr/mL (range 4.71-20.1 µg•hr/mL), respectively. In  
44 patients with respiratory and urinary tract infections (n=1423), similar estimates of  
45 systemic drug exposure were determined using a population pharmacokinetics analysis  
46 (geometric mean AUC (0-24), 8.36 µg•hr/mL; range 3.2 – 47.7 µg•hr/mL).

47 The pharmacokinetics of gemifloxacin were not significantly altered when a 320  
48 mg dose was administered with a high-fat meal. Therefore FACTIVE tablets may be  
49 administered without regard to meals.

50

### 51 **Distribution**

52 *In vitro* binding of gemifloxacin to plasma proteins in healthy subjects is approximately  
53 60 to 70% and is concentration independent. After repeated doses, the *in vivo* plasma  
54 protein binding in healthy elderly and young subjects ranged from 55% to 73% and was  
55 unaffected by age. Renal impairment does not significantly affect the protein binding of  
56 gemifloxacin. The blood-to-plasma concentration ratio of gemifloxacin was 1.2:1. The  
57 geometric mean for V<sub>dss</sub>/F is 4.18 L/kg (range, 1.66 – 12.12 L/kg).

58 Gemifloxacin is widely distributed throughout the body after oral administration.  
59 Concentrations of gemifloxacin in bronchoalveolar lavage fluid exceed those in the  
60 plasma. Gemifloxacin penetrates well into lung tissue and fluids. After five daily doses  
61 of 320 mg gemifloxacin, concentrations in plasma, bronchoalveolar macrophages,  
62 epithelial lining fluid and bronchial mucosa at approximately 2 hours were as in Table 1:  
63

64

**Table 1. Gemifloxacin Concentrations in Plasma and Tissues (320 mg Oral Dosing)**

Tissue	Concentration (mean ± SD)	Ratio compared with plasma (mean±SD)
Plasma	1.40 (0.442) µg/mL	---
Bronchoalveolar Macrophages	107 (77) µg/g	90.5 (106.3)
Epithelial Lining Fluid	2.69 (1.96) µg/mL	1.99 (1.32)
Bronchial Mucosa	9.52 (5.15) µg/g	7.21 (4.03)

65

### 66 **Metabolism**

67 Gemifloxacin is metabolized to a limited extent by the liver. The unchanged compound  
68 is the predominant drug-related component detected in plasma (approximately 65%) up  
69 to 4 hours after dosing. All metabolites formed are minor (<10% of the administered oral  
70 dose); the principal ones are N-acetyl gemifloxacin, the E-isomer of gemifloxacin and the  
71 carbamyl glucuronide of gemifloxacin. Cytochrome P450 enzymes do not play an  
72 important role in gemifloxacin metabolism, and the metabolic activity of these enzymes  
73 is not significantly inhibited by gemifloxacin.

74

### 75 **Excretion**

76 Gemifloxacin and its metabolites are excreted via dual routes of excretion. Following  
77 oral administration of gemifloxacin to healthy subjects, a mean (± SD) of 61 ± 9.5% of  
78 the dose was excreted in the feces and 36 ± 9.3% in the urine as unchanged drug and  
79 metabolites. The mean (± SD) renal clearance following repeat doses of 320 mg was  
80 approximately 11.6 ± 3.9 L/hr (range 4.6-17.6 L/hr), which indicates active secretion is  
81 involved in the renal excretion of gemifloxacin. The mean (± SD) plasma elimination

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82 half-life at steady state following 320 mg to healthy subjects was approximately  $7 \pm 2$   
83 hours (range 4-12 hours).

84

85 ***Special Populations***

86 **Pediatric:** The pharmacokinetics of gemifloxacin in pediatric subjects have not been  
87 studied.

88

89 **Geriatric:** In adult subjects, the pharmacokinetics of gemifloxacin are not affected by  
90 age.

91

92 **Gender:** There are no significant differences between gemifloxacin pharmacokinetics in  
93 males and females when differences in body weight are taken into account. Population  
94 pharmacokinetic studies indicated that following administration of 320 mg gemifloxacin,  
95 AUC values were approximately 10% higher in healthy female patients compared to  
96 males. Males and females had mean AUC values of 7.98  $\mu\text{g}\cdot\text{hr}/\text{mL}$  (range, 3.21 – 42.71  
97  $\mu\text{g}\cdot\text{hr}/\text{mL}$ ) and 8.80  $\mu\text{g}\cdot\text{hr}/\text{mL}$  (range, 3.33 – 47.73  $\mu\text{g}\cdot\text{hr}/\text{mL}$ ), respectively. No  
98 gemifloxacin dosage adjustment based on gender is necessary.

99

100 **Hepatic Insufficiency:** The pharmacokinetics following a single 320 mg dose of  
101 gemifloxacin were studied in patients with mild (Child-Pugh Class A) to moderate  
102 (Child-Pugh Class B) liver disease. There was a mean increase in AUC (0-inf) of 34%  
103 and a mean increase in C<sub>max</sub> of 25% in these patients with hepatic impairment compared  
104 to healthy volunteers.

105 The pharmacokinetics of a single 320 mg dose of gemifloxacin were also studied  
106 in patients with severe hepatic impairment (Child-Pugh Class C). There was a mean  
107 increase in AUC (0-inf) of 45% and a mean increase in C<sub>max</sub> of 41% in these subjects  
108 with hepatic impairment compared to healthy volunteers.

109 These average pharmacokinetic increases are not considered to be clinically  
110 significant. There was no significant change in plasma elimination half-life in the mild,  
111 moderate or severe hepatic impairment patients. No dosage adjustment is recommended  
112 in patients with mild (Child-Pugh Class A), moderate (Child-Pugh Class B) or severe  
113 (Child-Pugh Class C) hepatic impairment. (See **DOSAGE AND ADMINISTRATION.**)

114

115 **Renal Insufficiency:** Results from population pharmacokinetic and clinical  
116 pharmacology studies with repeated 320 mg doses indicate the clearance of gemifloxacin  
117 is reduced and the plasma elimination is prolonged, leading to an average increase in  
118 AUC values of approximately 70% in patients with renal insufficiency. In the  
119 pharmacokinetic studies, gemifloxacin C<sub>max</sub> was not significantly altered in subjects  
120 with renal insufficiency. Dose adjustment in patients with creatinine clearance >40  
121 mL/min is not required. Modification of the dosage is recommended for patients with  
122 creatinine clearance  $\leq 40$  mL/min. (See **DOSAGE AND ADMINISTRATION.**)

123

124 Hemodialysis removes approximately 20 to 30% of an oral dose of gemifloxacin from  
125 plasma.

126

127

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128 ***Photosensitivity Potential***

129 In a study of the skin response to ultraviolet and visible radiation conducted in 40 healthy  
130 volunteers, the minimum erythematous dose (MED) was assessed following  
131 administration of either gemifloxacin 160 mg once daily, gemifloxacin 320 mg once  
132 daily, ciprofloxacin 500 mg BID, or placebo for 7 days. At 5 of the 6 wavelengths tested  
133 (295-430 nm), the photosensitivity potential of gemifloxacin was not statistically  
134 different from placebo. At 365 nm (UVA region), gemifloxacin showed a  
135 photosensitivity potential similar to that of ciprofloxacin 500 mg BID and the  
136 photosensitivity potential for both drugs were statistically greater than that of placebo.  
137 Photosensitivity reactions were reported rarely in clinical trials with gemifloxacin  
138 (0.039%). (See **ADVERSE REACTIONS**.)

139

140 ***Drug-Drug Interactions***

141 Antacids/Di- and Trivalent Cations: The systemic availability of gemifloxacin is  
142 significantly reduced when an aluminum- and magnesium- containing antacid is  
143 concomitantly administered (AUC decreased 85%; Cmax decreased 87%).

144 Administration of an aluminum- and magnesium- containing antacid or ferrous sulfate  
145 (325 mg) at 3 hours before or at 2 hours after gemifloxacin did not significantly alter the  
146 systemic availability of gemifloxacin. Therefore, aluminum- and/or magnesium-  
147 containing antacids, ferrous sulfate (iron), multivitamin preparations containing zinc or  
148 other metal cations, or Videx® (didanosine) chewable/buffered tablets or the pediatric  
149 powder for oral solution should not be taken within 3 hours before or 2 hours after taking  
150 FACTIVE tablets.

151 Calcium carbonate (1000 mg) given either 2 hr before or 2 hr after gemifloxacin  
152 administration showed no notable reduction in gemifloxacin systemic availability.  
153 Calcium carbonate administered simultaneously with gemifloxacin resulted in a small,  
154 not clinically significant, decrease in gemifloxacin exposure [AUC (0-inf) decreased 21%  
155 and Cmax decreased].

156

157 Sucralfate: When sucralfate (2 g) was administered 3 hours prior to gemifloxacin, the oral  
158 bioavailability of gemifloxacin was significantly reduced (53% decrease in AUC; 69%  
159 decrease in Cmax). When sucralfate (2 g) was administered 2 hours after gemifloxacin,  
160 the oral bioavailability of gemifloxacin was not significantly affected; therefore  
161 FACTIVE should be taken at least 2 hours before sucralfate. (See **PRECAUTIONS**.)

162

163 In Vitro Metabolism: Results of *in vitro* inhibition studies indicate that hepatic  
164 cytochrome P450 (CYP450) enzymes do not play an important role in gemifloxacin  
165 metabolism. Therefore gemifloxacin should not cause significant *in vivo* pharmacokinetic  
166 interactions with other drugs that are metabolized by CYP450 enzymes.

167

168 Theophylline: Gemifloxacin 320 mg at steady-state did not affect the repeat dose  
169 pharmacokinetics of theophylline (300 to 400 mg BID to healthy male subjects).

170

171 Digoxin: Gemifloxacin 320 mg at steady-state did not affect the repeat dose  
172 pharmacokinetics of digoxin (0.25 mg once daily to healthy elderly subjects).

173

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174 Oral Contraceptives: The effect of an oral estrogen/progesterone contraceptive product  
175 (once daily for 21 days) on the pharmacokinetics of gemifloxacin (320 mg once daily for  
176 6 days) in healthy female subjects indicates that concomitant administration caused an  
177 average reduction in gemifloxacin AUC and C<sub>max</sub> of 19% and 12%. These changes are  
178 not considered clinically significant. Gemifloxacin 320 mg at steady-state did not affect  
179 the repeat dose pharmacokinetics of an ethinylestradiol/levonorgestrol oral contraceptive  
180 product (30 µg/150 µg once daily for 21 days to healthy female subjects).

181

182 Cimetidine: Co-administration of a single dose of 320 mg gemifloxacin with cimetidine  
183 400 mg four times daily for 7 days resulted in slight average increases in gemifloxacin  
184 AUC(0-inf) and C<sub>max</sub> of 10% and 6%, respectively. These increases are not considered  
185 clinically significant.

186

187 Omeprazole: Co-administration of a single dose of 320 mg gemifloxacin with  
188 omeprazole 40 mg once daily for 4 days resulted in slight average increases in  
189 gemifloxacin AUC(0-inf) and C<sub>max</sub> of 10% and 11%, respectively. These increases are  
190 not considered clinically significant.

191

192 Warfarin: Administration of repeated doses of gemifloxacin (320 mg once daily for 7  
193 days) to healthy subjects on stable warfarin therapy had no significant effect on warfarin-  
194 induced anticoagulant activity (i.e., International Normalized Ratios for Prothrombin  
195 Time). (See **PRECAUTIONS: Drug Interactions**.)

196

197 Probenecid: Administration of a single dose of 320 mg gemifloxacin to healthy subjects  
198 who also received repeat doses of probenecid (total dose = 4.5 g) reduced the mean renal  
199 clearance of gemifloxacin by approximately 50%, resulting in a mean increase of 45% in  
200 gemifloxacin AUC (0-inf) and a prolongation of mean half-life by 1.6 hours. Mean  
201 gemifloxacin C<sub>max</sub> increased 8%.

202

## 203 **MICROBIOLOGY**

204 Gemifloxacin has *in vitro* activity against a wide range of Gram-negative and Gram-  
205 positive microorganisms. Gemifloxacin is bactericidal with minimum bactericidal  
206 concentrations (MBCs) generally within one dilution of the minimum inhibitory  
207 concentrations (MICs). Gemifloxacin acts by inhibiting DNA synthesis through the  
208 inhibition of both DNA gyrase and topoisomerase IV (TOPO IV), which are essential for  
209 bacterial growth. *Streptococcus pneumoniae* showing mutations in both DNA gyrase and  
210 TOPO IV (double mutants) are resistant to most fluoroquinolones. Gemifloxacin has the  
211 ability to inhibit both enzyme systems at therapeutically relevant drug levels in *S.*  
212 *pneumoniae* (dual targeting), and has MIC values that are still in the susceptible range for  
213 some of these double mutants. However, the presence of double mutants was not  
214 evaluated in clinical trials; therefore, the clinical significance of these *in vitro* data are  
215 unknown.

216 The mechanism of action of quinolones, including gemifloxacin, is different from  
217 that of macrolides, beta-lactams, aminoglycosides, or tetracyclines; therefore,  
218 microorganisms resistant to these classes of drugs may be susceptible to gemifloxacin

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219 and other quinolones. There is no known cross-resistance between gemifloxacin and the  
220 above mentioned classes of antimicrobials.

221 The main mechanism of fluoroquinolone resistance is due to mutations in DNA  
222 gyrase and/or TOPO IV. Resistance to gemifloxacin develops slowly via multistep  
223 mutations and efflux in a manner similar to other fluoroquinolones. The frequency of  
224 spontaneous mutation is low ( $10^{-7}$  to  $<10^{-10}$ ). Although cross-resistance has been  
225 observed between gemifloxacin and other fluoroquinolones, some microorganisms  
226 resistant to other fluoroquinolones may be susceptible to gemifloxacin.

227 Gemifloxacin has been shown to be active against most strains of the following  
228 microorganisms, both *in vitro* and in clinical infections as described in the  
229 **INDICATIONS AND USAGE** section.

230

231 **Aerobic Gram-positive microorganisms**

232 *Streptococcus pneumoniae* (including multi-drug resistant strains [MDRSP])\*

233 \*MDRSP, multi-drug resistant *Streptococcus pneumoniae*, includes isolates previously  
234 known as PRSP (penicillin-resistant *Streptococcus pneumoniae*), and are strains resistant  
235 to two or more of the following antibiotics: penicillin (MIC  $\geq 2$   $\mu\text{g/mL}$ ), 2nd generation  
236 cephalosporins (e.g., cefuroxime), macrolides, tetracyclines and  
237 trimethoprim/sulfamethoxazole.

238

239 **Aerobic Gram-negative microorganisms**

240 *Haemophilus influenzae*

241 *Haemophilus parainfluenzae*

242 *Klebsiella pneumoniae* (many strains are only moderately susceptible)

243 *Moraxella catarrhalis*

244

245 **Other microorganisms**

246 *Chlamydia pneumoniae*

247 *Mycoplasma pneumoniae*

248

249 The following data are available, **but their clinical significance is unknown.**

250

251 Gemifloxacin exhibits *in vitro* minimal inhibitory concentrations (MICs) of 0.25  
252  $\mu\text{g/mL}$  or less against most ( $\geq 90\%$ ) strains of the following microorganisms; however,  
253 the safety and effectiveness of gemifloxacin in treating clinical infections due to these  
254 microorganisms has not been established in adequate and well-controlled clinical trials:

255

256 **Aerobic Gram-positive microorganisms**

257 *Staphylococcus aureus* (methicillin-susceptible strains only)

258 *Streptococcus pyogenes*

259

260 **Aerobic Gram-negative microorganisms**

261 *Acinetobacter lwoffii*

262 *Klebsiella oxytoca*

263 *Legionella pneumophila*

264 *Proteus vulgaris*

---

265 **Susceptibility Tests**

266 **Dilution techniques:** Quantitative methods are used to determine antimicrobial  
267 minimum inhibitory concentrations (MICs). These MICs provide estimates of the  
268 susceptibility of bacteria to antimicrobial compounds. The MICs should be determined  
269 using a standardized procedure. Standardized procedures are based on a dilution method<sup>1</sup>  
270 (broth or agar) or equivalent with standardized inoculum concentrations and standardized  
271 concentrations of gemifloxacin powder. The MICs should be interpreted according to the  
272 following criteria:

273

274 For testing *Klebsiella pneumoniae*:

<u>MIC (µg/mL)</u>	<u>Interpretation</u>
275 ≤0.25	Susceptible (S)
276 0.5	Intermediate (I)
277 ≥1.0	Resistant (R)

278

279 For testing *Haemophilus influenzae* and *Haemophilus parainfluenzae*<sup>a</sup>:

<u>MIC (µg/mL)</u>	<u>Interpretation</u>
280 ≤0.12	Susceptible (S)

281

282 <sup>a</sup> This interpretive standard is applicable only to broth microdilution susceptibility testing  
283 with *Haemophilus influenzae* and *Haemophilus parainfluenzae* using *Haemophilus* Test  
284 Medium (HTM)<sup>1</sup>.

285

286 The current absence of data on resistant strains precludes defining any results  
287 other than “Susceptible”. Strains yielding MIC results suggestive of a “nonsusceptible”  
288 category should be submitted to a reference laboratory for further testing.

289

290 For testing *Streptococcus pneumoniae*<sup>b</sup>:

<u>MIC (µg/mL)</u>	<u>Interpretation</u>
291 ≤0.12	Susceptible (S)
292 0.25	Intermediate (I)
293 ≥0.5	Resistant (R)

294

295 <sup>b</sup> These interpretive standards are applicable only to broth microdilution susceptibility  
296 tests using cation-adjusted Mueller-Hinton broth with 2-5% lysed horse blood.

297

298 A report of “Susceptible” indicates that the pathogen is likely to be inhibited if the  
299 antimicrobial compound in the blood reaches the concentration usually achievable. A  
300 report of “Intermediate” indicates that the result should be considered equivocal, and if  
301 the microorganism is not fully susceptible to alternative, clinically feasible drugs, the test  
302 should be repeated. This category implies possible clinical applicability in body sites  
303 where the drug is physiologically concentrated or in situations where high dosage of drug  
304 can be used. This category also provides a buffer zone, which prevents small  
305 uncontrolled technical factors from causing major discrepancies in interpretation. A  
306 report of “Resistant” indicates that the pathogen is not likely to be inhibited if the

307

310 antimicrobial compound in the blood reaches the concentration usually achievable; other  
311 therapy should be selected.

312 Standardized susceptibility test procedures require the use of laboratory control  
313 microorganisms to control the technical aspects of the laboratory procedures. Standard  
314 gemifloxacin powder should provide the following MIC values:

315

<u>Microorganism</u>		<u>MIC Range (µg/mL)</u>
316 <i>Enterococcus faecalis</i>	ATCC 29212	0.016-0.12
317 <i>Escherichia coli</i>	ATCC 25922	0.004-0.016
318 <i>Haemophilus influenzae</i>	ATCC 49247	0.002-0.008 <sup>c</sup>
319 <i>Streptococcus pneumoniae</i>	ATCC 49619	0.008-0.03 <sup>d</sup>

320  
321  
322 <sup>c</sup> This quality control range is applicable to only *H. influenzae* ATCC 49247 tested by a  
323 broth microdilution procedure using *Haemophilus* Test Medium (HTM)<sup>1</sup>.

324  
325 <sup>d</sup> This quality control range is applicable to only *S. pneumoniae* ATCC 49619 tested by a  
326 broth microdilution procedure using cation-adjusted Mueller-Hinton broth with 2-5%  
327 lysed horse blood.

328  
329 **Diffusion Techniques:** Quantitative methods that require measurement of zone  
330 diameters also provide reproducible estimates of the susceptibility of bacteria to  
331 antimicrobial compounds. One such standardized procedure<sup>2</sup> requires the use of  
332 standardized inoculum concentrations. This procedure uses paper disks impregnated with  
333 5 µg gemifloxacin to test the susceptibility of microorganisms to gemifloxacin.

334 Reports from the laboratory providing results of the standard single-disk  
335 susceptibility test with a 5 µg gemifloxacin disk should be interpreted according to the  
336 following criteria:

337  
338 For testing *Klebsiella pneumoniae*:

<u>Zone Diameter (mm)</u>	<u>Interpretation</u>
339 ≥20	Susceptible (S)
340 16-19	Intermediate (I)
341 ≤15	Resistant (R)

342  
343  
344 For testing *Haemophilus influenzae* and *Haemophilus parainfluenzae*<sup>e</sup>:

<u>Zone Diameter (mm)</u>	<u>Interpretation</u>
345 ≥18	Susceptible (S)

346  
347  
348 <sup>e</sup> This interpretive standard is applicable only to disk diffusion susceptibility testing with  
349 *Haemophilus influenzae* and *Haemophilus parainfluenzae* using *Haemophilus* Test  
350 Medium (HTM).<sup>2</sup>

351  
352 The current absence of data on resistant strains precludes defining any results  
353 other than “Susceptible”. Strains yielding zone diameter results suggestive of a  
354 “nonsusceptible” category should be submitted to a reference laboratory for further  
355 testing.



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356 For testing *Streptococcus pneumoniae*<sup>f</sup>:

357	<u>Zone Diameter (mm)</u>	<u>Interpretation</u>
358	≥23	Susceptible (S)
359	20-22	Intermediate (I)
360	≤19	Resistant (R)

361

362 <sup>f</sup> These zone diameter standards apply only to tests performed using Mueller-Hinton agar  
363 supplemented with 5% defibrinated sheep blood incubated in 5% CO<sub>2</sub>.

364

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

366 Interpretation involves correlation of the diameter obtained in the disk test with the MIC  
367 for gemifloxacin.

368

369 As with standardized dilution techniques, diffusion methods require the use of  
370 laboratory control microorganisms that are used to control the technical aspects of the  
371 laboratory procedures. For the diffusion technique, the 5 µg gemifloxacin disk should  
372 provide the following zone diameters in these laboratory quality control strains:

373

374	<u>Microorganism</u>	<u>Zone Diameter (mm)</u>
375	<i>Escherichia coli</i> ATCC 25922	29-36
376	<i>Haemophilus influenzae</i> ATCC 49247	30-37 <sup>g</sup>
377	<i>Streptococcus pneumoniae</i> ATCC 49619	28-34 <sup>h</sup>

378

379 <sup>g</sup> This quality control range is applicable to only *H. influenzae* ATCC 49247 tested by a  
380 disk diffusion procedure using *Haemophilus* Test Medium (HTM)<sup>2</sup>.

381

382 <sup>h</sup> This quality control range is applicable to only *S. pneumoniae* ATCC 49619 tested by a  
383 disk diffusion procedure using Mueller-Hinton agar supplemented with 5% defibrinated  
384 sheep blood and incubated in 5% CO<sub>2</sub>.

385

## 386 INDICATIONS AND USAGE

387 FACTIVE is indicated for the treatment of infections caused by susceptible strains of the  
388 designated microorganisms in the conditions listed below. (See **DOSAGE AND**  
389 **ADMINISTRATION** and **CLINICAL STUDIES**.)

390

391 **Acute bacterial exacerbation of chronic bronchitis** caused by *Streptococcus*  
392 *pneumoniae*, *Haemophilus influenzae*, *Haemophilus parainfluenzae*, or *Moraxella*  
393 *catarrhalis*.

394

395 **Community-acquired pneumonia** (of mild to moderate severity) caused by  
396 *Streptococcus pneumoniae* (including multi-drug resistant strains [MDRSP])\*  
397 *Haemophilus influenzae*, *Moraxella catarrhalis*, *Mycoplasma pneumoniae*, *Chlamydia*  
398 *pneumoniae*, or *Klebsiella pneumoniae*.

399

400 \*MDRSP, multi-drug resistant *Streptococcus pneumoniae*, includes isolates  
401 previously known as PRSP (penicillin-resistant *Streptococcus pneumoniae*), and are

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402 strains resistant to two or more of the following antibiotics: penicillin (MIC  $\geq 2$   $\mu\text{g/mL}$ ),  
403 2nd generation cephalosporins (e.g., cefuroxime), macrolides, tetracyclines and  
404 trimethoprim/sulfamethoxazole.

405

406 To reduce the development of drug-resistant bacteria and maintain the  
407 effectiveness of FACTIVE and other antibacterial drugs, FACTIVE should be used only  
408 to treat infections that are proven or strongly suspected to be caused by susceptible  
409 bacteria. When culture and susceptibility information are available, they should be  
410 considered in selecting or modifying antibacterial therapy. In the absence of such data,  
411 local epidemiology and susceptibility patterns may contribute to the empiric selection of  
412 therapy.

413

#### 414 **CONTRAINDICATIONS**

415 FACTIVE is contraindicated in patients with a history of hypersensitivity to  
416 gemifloxacin, fluoroquinolone antibiotic agents, or any of the product components.

417

#### 418 **WARNINGS**

419 **THE SAFETY AND EFFECTIVENESS OF FACTIVE IN CHILDREN,**  
420 **ADOLESCENTS (LESS THAN 18 YEARS OF AGE), PREGNANT WOMEN,**  
421 **AND LACTATING WOMEN HAVE NOT BEEN ESTABLISHED. (See**  
422 **PRECAUTIONS: Pediatric Use, Pregnancy and Nursing Mothers subsections.)**

423

424 ***QT Effects:*** Fluoroquinolones may prolong the QT interval in some patients. FACTIVE  
425 should be avoided in patients with a history of prolongation of the QTc interval, patients  
426 with uncorrected electrolyte disorders (hypokalemia or hypomagnesemia), and patients  
427 receiving Class IA (e.g., quinidine, procainamide) or Class III (e.g., amiodarone, sotalol)  
428 antiarrhythmic agents.

429

430 Pharmacokinetic studies between gemifloxacin and drugs that prolong the QTc  
431 interval such as erythromycin, antipsychotics, and tricyclic antidepressants have not been  
432 performed. FACTIVE should be used with caution when given concurrently with these  
433 drugs, as well as in patients with ongoing proarrhythmic conditions, such as clinically  
434 significant bradycardia or acute myocardial ischemia. No cardiovascular morbidity or  
435 mortality attributable to QTc prolongation occurred with FACTIVE treatment in over  
436 8119 patients, including 707 patients concurrently receiving drugs known to prolong the  
437 QTc interval and 7 patients with hypokalemia.

438

439 The likelihood of QTc prolongation may increase with increasing dose of the  
440 drug; therefore, the recommended dose should not be exceeded especially in patients with  
441 renal or hepatic impairment where the C<sub>max</sub> and AUC are slightly higher. QTc  
442 prolongation may lead to an increased risk for ventricular arrhythmias including torsades  
443 de pointes. The maximal change in the QTc interval occurs approximately 5-10 hours  
444 following oral administration of gemifloxacin.

445

446 ***Hypersensitivity Reactions:*** Serious hypersensitivity and/or anaphylactic reactions have  
447 been reported in patients receiving fluoroquinolone therapy, including FACTIVE.

448

449 Hypersensitivity reactions reported in patients receiving fluoroquinolone therapy have  
450 occasionally been fatal. These reactions may occur following the first dose. Some

---

448 reactions have been accompanied by cardiovascular collapse, hypotension/shock, seizure,  
449 loss of consciousness, tingling, angioedema (including tongue, laryngeal, throat or facial  
450 edema/swelling), airway obstruction (including bronchospasm, shortness of breath and  
451 acute respiratory distress), dyspnea, urticaria, itching and other serious skin reactions.

452 FACTIVE should be discontinued immediately at the appearance of any sign of  
453 an immediate type I hypersensitivity skin rash or any other manifestation of a  
454 hypersensitivity reaction; the need for continued fluoroquinolone therapy should be  
455 evaluated. As with other drugs, serious acute hypersensitivity reactions may require  
456 treatment with epinephrine and other resuscitative measures, including oxygen,  
457 intravenous fluids, antihistamines, corticosteroids, pressor amines and airway  
458 management as clinically indicated. (See **PRECAUTIONS** and **ADVERSE**  
459 **REACTIONS**.)

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

- 465 • fever, rash or severe dermatologic reactions (e.g., toxic epidermal necrolysis,  
466 Stevens-Johnson Syndrome);
- 467 • vasculitis; arthralgia; myalgia; serum sickness;
- 468 • allergic pneumonitis;
- 469 • interstitial nephritis; acute renal insufficiency or failure;
- 470 • hepatitis; jaundice; acute hepatic necrosis or failure;
- 471 • anemia, including hemolytic and aplastic;
- 472 • thrombocytopenia, including thrombotic thrombocytopenic purpura; leukopenia  
473 agranulocytosis; pancytopenia; and/or other hematologic abnormalities.

474  
475 The drug should be discontinued immediately at the first appearance of a skin  
476 rash, jaundice, or any other sign of hypersensitivity and supportive measures instituted  
477 (See **PRECAUTIONS: Information for Patients** and **ADVERSE REACTIONS**).

478  
479 **Peripheral Neuropathy:** Rare cases of sensory or sensorimotor axonal polyneuropathy  
480 affecting small and/or large axons resulting in paresthesias, hypoesthesias, dysesthesias  
481 and weakness have been reported in patients receiving quinolones.

482  
483 **Tendon Effects:** Ruptures of the shoulder, hand, Achilles tendon or other tendons that  
484 required surgical repair or resulted in prolonged disability have been reported in patients  
485 receiving quinolones. Post-marketing surveillance reports indicate that this risk may be  
486 increased in patients receiving concomitant corticosteroids, especially the elderly.  
487 FACTIVE should be discontinued if the patient experiences pain, inflammation, or  
488 rupture of a tendon. Patients should rest and refrain from exercise until the diagnosis of  
489 tendonitis or tendon rupture has been excluded. Tendon rupture can occur during or after  
490 therapy with quinolones.

491  
492 **CNS Effects:** In clinical studies with FACTIVE, central nervous system (CNS) effects  
493 have been reported infrequently. As with other fluoroquinolones, FACTIVE should be

494 used with caution in patients with CNS diseases such as epilepsy or patients predisposed  
495 to convulsions. Although not seen in FACTIVE clinical trials, convulsions, increased  
496 intracranial pressure, and toxic psychosis have been reported in patients receiving other  
497 fluoroquinolones. CNS stimulation which may lead to tremors, restlessness, anxiety,  
498 lightheadedness, confusion, hallucinations, paranoia, depression, insomnia, and rarely  
499 suicidal thoughts or acts may also be caused by other fluoroquinolones. If these reactions  
500 occur in patients receiving FACTIVE, the drug should be discontinued and appropriate  
501 measures instituted.

502

503 ***Clostridium difficile* Associated Diarrhea:** *Clostridium difficile* associated diarrhea  
504 (CDAD) has been reported with use of nearly all antibacterial agents, including  
505 FACTIVE, and may range in severity from mild diarrhea to fatal colitis. Treatment with  
506 antibacterial agents alters the normal flora of the colon leading to overgrowth of *C.*  
507 *difficile*.

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

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

518

## 519 PRECAUTIONS

520 **General:** Prescribing FACTIVE in the absence of a proven or strongly suspected  
521 bacterial infection is unlikely to provide benefit to the patient and increases the risk of the  
522 development of drug-resistant bacteria.

523

524 **Rash:** In clinical studies, rash occurred more often with FACTIVE than with therapy  
525 with comparator agents (2.7% vs 0.6%). Increasing incidence of rash was associated  
526 with younger age (especially below 40), female gender, use of hormone replacement  
527 therapy and longer durations of therapy (see Table 2). Urticarial reactions, some of  
528 which were not classified as rash, were more common in FACTIVE patients than in  
529 comparator patients (0.6% vs 0.2%). FACTIVE should be discontinued in patients  
530 developing a rash or urticaria while on treatment. (See ADVERSE REACTIONS and  
531 CLINICAL STUDIES.)

532

533 **Table 2. Rash Incidence in FACTIVE Treated Patients from the Clinical Studies**  
534 **Population\* by Gender, Age, and Duration of Therapy**

Gender & Age (yr) Category	Duration of FACTIVE Therapy			
	5 days	7 days	10 days**	14 days**
Female < 40	10/399 (2.5%)	49/407 (12.0%)	20/131 (15.3%)	7/31 (22.6%)
Female ≥ 40	30/1438 (2.1%)	34/887 (3.8%)	19/308 (6.2%)	10/126 (7.9%)
Male < 40	6/356 (1.7%)	26/453 (5.7%)	7/74 (9.5%)	3/39 (7.7%)

---

<b>Male ≥ 40</b>	10/1503 (0.7%)	26/984 (2.6%)	9/345 (2.6%)	3/116 (2.6%)
<b>Totals</b>	56/3696 (1.5%)	135/2732 (4.9%)	55/858 (6.4%)	23/312 (7.4%)

535 \*includes patients from studies of community-acquired pneumonia, acute bacterial  
 536 exacerbation of chronic bronchitis, and other indications

537 \*\*exceeds the recommended duration of therapy (see **DOSAGE AND**  
 538 **ADMINISTRATION**)

539

540 The most common form of rash associated with FACTIVE was described as  
 541 maculopapular and mild to moderate in severity. Eighty percent of rashes resolved  
 542 within 14 days. Approximately 10% of the rashes (0.5% of all patients) were described  
 543 as of severe intensity and approximately 10% percent of those with rash were treated with  
 544 systemic steroids. There were no documented cases in the clinical trials of more serious  
 545 skin reactions known to be associated with significant morbidity or mortality.

546

547 Photosensitivity reactions have been reported rarely in clinical trials with  
 548 FACTIVE. (See **CLINICAL PHARMACOLOGY**.) However, as with all drugs of this  
 549 class, it is recommended that patients avoid unnecessary exposure to strong sunlight or  
 550 artificial UV rays (e.g., sunlamps, solariums), and should be advised of the appropriate  
 551 use of broad spectrum sun block if in bright sunlight. Treatment should be discontinued  
 552 if a photosensitivity reaction is suspected.

553

554 **Hepatic Effects:** Liver enzyme elevations (increased ALT and/or AST) occurred at  
 555 similar rates in patients receiving FACTIVE 320 mg daily relative to comparator  
 556 antimicrobial agents (ciprofloxacin, levofloxacin, clarithromycin/cefuroxime axetil,  
 557 amoxicillin/clavulanate potassium, and ofloxacin). In patients who received  
 558 gemifloxacin at doses of 480 mg per day or greater there was an increased incidence of  
 559 elevations in liver enzymes. (See **ADVERSE REACTIONS**.)

560

561 There were no clinical symptoms associated with these liver enzyme elevations.  
 562 The liver enzyme elevations resolved following cessation of therapy. The recommended  
 563 dose of FACTIVE 320 mg daily should not be exceeded and the recommended length of  
 564 therapy should not be exceeded. (See **DOSAGE AND ADMINISTRATION**.)

564

565 **Renal Effects:** Alteration of the dosage regimen is necessary for patients with  
 566 impairment of renal function (creatinine clearance ≤40 mL/min). (See **DOSAGE AND**  
 567 **ADMINISTRATION**.)

568

569 Adequate hydration of patients receiving FACTIVE should be maintained to  
 570 prevent the formation of a highly concentrated urine.

570

571 **Information for Patients**

572 Patients should be counseled:

573

- 574 • that antibacterial drugs including FACTIVE should only be used to treat bacterial  
 575 infections. They do not treat viral infections (e.g., the common cold). When  
 576 FACTIVE is prescribed to treat a bacterial infection, patients should be told that  
 577 although it is common to feel better early in the course of therapy, the medication  
 578 should be taken exactly as directed. Skipping doses or not completing the full course  
 of therapy may (1) decrease effectiveness of the immediate treatment and (2) increase

---

579 the likelihood that bacteria will develop resistance and will not be treatable by  
580 FACTIVE or other antibacterial drugs in the future;

- 581 • that FACTIVE has been associated with rash and hives. Rash occurs more commonly  
582 in those under 40, especially women and in women on hormone replacement therapy.  
583 The incidence of rash increases with duration more than 5 days and particularly  
584 longer than 7 days. Patients should discontinue FACTIVE and call their healthcare  
585 provider if they develop a rash;
- 586 • that FACTIVE may be associated with hypersensitivity reactions, including  
587 anaphylactic reactions, even following a single dose; patients should immediately  
588 discontinue the drug at the sign of a rash or other allergic reaction and seek medical  
589 care;
- 590 • that diarrhea is a common problem caused by antibiotics which usually ends when the  
591 antibiotic is discontinued. Sometimes after starting treatment with antibiotics,  
592 patients can develop watery and bloody stools (with or without stomach cramps and  
593 fever) even as late as two or more months after having taken the last dose of the  
594 antibiotic. If this occurs, patients should contact their physician as soon as possible;
- 595 • that FACTIVE may cause changes in the electrocardiogram (QTc interval  
596 prolongation);
- 597 • that FACTIVE should be avoided in patients receiving Class IA (e.g., quinidine,  
598 procainamide) or Class III (e.g., amiodarone, sotalol) antiarrhythmic agents;
- 599 • that FACTIVE should be used with caution in patients receiving drugs that affect the  
600 QTc interval such as cisapride, erythromycin, antipsychotics, and tricyclic  
601 antidepressants;
- 602 • to inform their physician of any personal or family history of QTc prolongation or  
603 proarrhythmic conditions such as hypokalemia, bradycardia, or recent myocardial  
604 ischemia;
- 605 • to contact their physician if they experience palpitations or fainting spells while  
606 taking FACTIVE;
- 607 • that FACTIVE may cause dizziness; if this occurs, patients should not operate an  
608 automobile or machinery or engage in activities requiring mental alertness or  
609 coordination;
- 610 • that they should discontinue FACTIVE therapy and inform their physician if they feel  
611 pain, tenderness or rupture of a tendon. Patients should rest and avoid exercise until  
612 the diagnosis of tendonitis or tendon rupture has been excluded. The risk of serious  
613 tendon disorders is higher in those over 65 years of age, especially those on steroids;
- 614 • that convulsions have been reported in patients receiving quinolones. Patients should  
615 notify their physician before taking FACTIVE if they have a history of convulsions,  
616 seizures, or epilepsy;
- 617 • that other central nervous system problems such as tremors, restlessness,  
618 lightheadedness, confusion and hallucinations may occur rarely;
- 619 • that phototoxicity has been reported with certain quinolones. The potential for  
620 FACTIVE to cause phototoxicity was low. In keeping with good clinical practice,  
621 avoid excessive sunlight or artificial ultraviolet light (e.g., tanning beds). If a  
622 sunburn-like reaction or skin eruption occurs, contact your physician; (See  
623 **CLINICAL PHARMACOLOGY: Photosensitivity Potential**);

- 
- 624 • that increases of the International Normalized Ratio (INR), or prothrombin time (PT),  
625 and/or clinical episodes of bleeding have been noted with concurrent administration  
626 of warfarin or its derivatives, and FACTIVE. Patients should notify their physicians  
627 if they are taking warfarin or its derivatives;
- 628 • to inform their physician of any other medications when taken concurrently with  
629 FACTIVE, including over-the-counter medications and dietary supplements;
- 630 • that FACTIVE may be taken with or without meals;
- 631 • to drink fluids liberally;
- 632 • not to take antacids containing magnesium and/or aluminum or products containing  
633 ferrous sulfate (iron), multivitamin preparations containing zinc or other metal  
634 cations, or Videx® (didanosine) chewable/buffered tablets or the pediatric powder for  
635 oral solution within 3 hours before or 2 hours after taking FACTIVE tablets;
- 636 • that FACTIVE should be taken at least 2 hours before sucralfate.

637

638 **Drug Interactions:** Administration of repeat doses of FACTIVE had no effect on the  
639 repeat dose pharmacokinetics of theophylline, digoxin or an

640 ethinylestradiol/levonorgestrol oral contraceptive product in healthy subjects. (See

641 **CLINICAL PHARMACOLOGY: Drug-Drug Interactions.**)

642 Concomitant administration of FACTIVE and calcium carbonate, cimetidine,  
643 omeprazole, or an estrogen/progesterone oral contraceptive produced minor changes in  
644 the pharmacokinetics of gemifloxacin, which were considered to be without clinical  
645 significance. (See **CLINICAL PHARMACOLOGY.**)

646 Concomitant administration of FACTIVE with probenecid resulted in a 45%  
647 increase in systemic exposure to gemifloxacin. (See **CLINICAL**  
648 **PHARMACOLOGY.**)

649 FACTIVE had no significant effect on the anticoagulant effect of warfarin in  
650 healthy subjects on stable warfarin therapy. However, post-marketing reports of  
651 increases in the INR, or PT, and/or clinical episodes of bleeding in patients have been  
652 noted with the use of quinolones, including FACTIVE, and warfarin, or its derivatives.  
653 In addition, infectious disease and its accompanying inflammatory process, age and  
654 general status of the patient are risk factors for increased anticoagulation activity.  
655 Therefore, the PT, INR or other suitable coagulation test should be closely monitored if a  
656 quinolone antimicrobial, including FACTIVE, is administered concomitantly with  
657 warfarin or its derivatives.

658 Quinolones form chelates with alkaline earth and transition metals. The  
659 absorption of oral gemifloxacin is significantly reduced by the concomitant  
660 administration of an antacid containing aluminum and magnesium. Magnesium- and/or  
661 aluminum-containing antacids, products containing ferrous sulfate (iron), multivitamin  
662 preparations containing zinc or other metal cations, or Videx® (didanosine)  
663 chewable/buffered tablets or the pediatric powder for oral solution should not be taken  
664 within 3 hours before or 2 hours after FACTIVE. Sucralfate should not be taken within 2  
665 hours of FACTIVE. (See **CLINICAL PHARMACOLOGY.**)

666

### 667 **Carcinogenesis, Mutagenesis, Impairment of Fertility**

668 *Carcinogenesis:* Long term studies in animals to determine the carcinogenic potential of  
669 gemifloxacin have not been conducted.

670

671 *Photocarcinogenesis:* Gemifloxacin did not shorten the time to development of UVR-  
672 induced skin tumors in hairless albino (Skh-1) mice; thus, it was not photocarcinogenic in  
673 this model. These mice received oral gemifloxacin and concurrent irradiation with  
674 simulated sunlight 5 days per week for 40 weeks followed by a 12-week treatment-free  
675 observation period. The daily dose of UV radiation used in this study was approximately  
676 1/3 of the minimal dose of UV radiation that would induce erythema in Caucasian  
677 humans. The median time to the development of skin tumors in the hairless mice was  
678 similar in the vehicle control group (36 weeks) and those given up to 100 mg/kg  
679 gemifloxacin daily (39 weeks). Following repeat doses of 100 mg/kg gemifloxacin per  
680 day, the mice had skin gemifloxacin concentrations of approximately 7.4 µg/g. Plasma  
681 levels following this dose were approximately 1.4 µg/mL in the mice around the time of  
682 irradiation. There are no data on gemifloxacin skin levels in humans, but the mouse  
683 plasma gemifloxacin levels are in the expected range of human plasma C<sub>max</sub> levels (0.7-  
684 2.6 µg/mL, with an overall mean of about 1.6 µg/mL) following multiple 320 mg oral  
685 doses.

686

687 *Mutagenesis:* Gemifloxacin was not mutagenic in 4 bacterial strains (TA 98, TA 100, TA  
688 1535, TA 1537) used in an Ames *Salmonella* reversion assay. It did not induce  
689 micronuclei in the bone marrow of mice following intraperitoneal doses of up to 40  
690 mg/kg and it did not induce unscheduled DNA synthesis in hepatocytes from rats which  
691 received oral doses of up to 1600 mg/kg. Gemifloxacin was clastogenic *in vitro* in the  
692 mouse lymphoma and human lymphocyte chromosome aberration assays. It was  
693 clastogenic *in vivo* in the rat micronucleus assay at oral and intravenous dose levels (≥800  
694 mg/kg and ≥40 mg/kg, respectively) that produced bone marrow toxicity.  
695 Fluoroquinolone clastogenicity is apparently due to inhibition of mammalian  
696 topoisomerase activity which has threshold implications.

697

698 *Impairment of Fertility:* Gemifloxacin did not affect the fertility of male or female rats at  
699 AUC levels following oral administration (216 and 600 mg/kg/day) that were  
700 approximately 3- to 4-fold higher than the AUC levels at the clinically recommended  
701 dose.

702

703 **Pregnancy: Teratogenic Effects. Pregnancy Category C.** Gemifloxacin treatment  
704 during organogenesis caused fetal growth retardation in mice (oral dosing at 450  
705 mg/kg/day), rats (oral dosing at 600 mg/kg/day) and rabbits (IV dosing at 40 mg/kg/day)  
706 at AUC levels which were 2-, 4- and 3-fold those in women given oral doses of 320 mg.  
707 In rats, this growth retardation appeared to be reversible in a pre- and postnatal  
708 development study (mice and rabbits were not studied for the reversibility of this effect).  
709 Treatment of pregnant rats at 8-fold clinical exposure (based upon AUC comparisons)  
710 caused fetal brain and ocular malformations in the presence of maternal toxicity. The  
711 overall no-effect exposure level in pregnant animals was approximately 0.8 to 3-fold  
712 clinical exposure.

713

714

The safety of FACTIVE in pregnant women has not been established. FACTIVE should not be used in pregnant women unless the potential benefit to the mother



---

715 outweighs the risk to the fetus. There are no adequate and well-controlled studies in  
716 pregnant women.

717

718 **Nursing Mothers:** Gemifloxacin is excreted in the breast milk of rats. There is no  
719 information on excretion of gemifloxacin into human milk. Therefore, FACTIVE should  
720 not be used in lactating women unless the potential benefit to the mother outweighs the  
721 risk.

722

723 **Pediatric Use:** Safety and effectiveness in children and adolescents less than 18 years of  
724 age have not been established. Fluoroquinolones, including gemifloxacin, cause  
725 arthropathy and osteochondrosis in immature animals. (See **WARNINGS**.)

726

727 **Geriatric Use:** Of the total number of subjects in clinical studies of FACTIVE, 29%  
728 (2314) were 65 and over, while 11% (865) were 75 and over. No overall difference in  
729 effectiveness was observed between these subjects and younger subjects; the adverse  
730 event rate for this group was similar to or lower than that for younger subjects with the  
731 exception that the incidence of rash was lower in geriatric patients compared to patients  
732 less than 40 years of age.

733 Elderly patients may be more susceptible to drug-associated effects on the QT  
734 interval. Therefore, FACTIVE should be avoided in patients taking drugs that can result  
735 in prolongation of the QT interval (e.g., Class IA or Class III antiarrhythmics) or in  
736 patients with risk factors for torsades de pointes (e.g., known QT prolongation,  
737 uncorrected hypokalemia).

738 Patients over 65 are at increased risk for developing severe tendon disorders  
739 including tendon rupture when being treated with a fluoroquinolone such as FACTIVE.  
740 This risk is further increased in patients receiving concomitant steroid therapy. Tendon  
741 rupture usually involves the Achilles, hand or shoulder tendons and can occur during  
742 therapy or up to a few months post completion of therapy. Caution should be used when  
743 prescribing FACTIVE to elderly patients especially those on corticosteroids. Patients  
744 should be informed of this potential side effect and advised to discontinue therapy and  
745 inform their physicians if any tendon symptoms occur.

746

## 747 **ADVERSE REACTIONS**

748 In clinical studies, 8119 patients received daily oral doses of 320 mg FACTIVE. In  
749 addition, 1797 healthy volunteers and 81 patients with renal or hepatic impairment  
750 received single or repeat doses of gemifloxacin in clinical pharmacology studies. The  
751 majority of adverse reactions experienced by patients in clinical trials were considered to  
752 be of mild to moderate severity.

753 FACTIVE was discontinued because of an adverse event (determined by the  
754 investigator to be possibly or probably related to drug) in 2.0% of patients, primarily due  
755 to rash (0.8%), nausea (0.3%), diarrhea (0.3%), urticaria (0.2%) and vomiting (0.2%).  
756 Comparator antibiotics were discontinued because of an adverse event at an overall  
757 comparable rate of 2.1%, primarily due to diarrhea (0.5%), nausea (0.4%), vomiting  
758 (0.3%), rash (0.3%), abdominal pain (0.2%), and vertigo (0.2%).

759 The most commonly reported adverse events with a frequency of  $\geq 2\%$  for patients  
760 receiving 320 mg of FACTIVE versus comparator drug (beta-lactam antibiotics),

761 macrolides or other fluoroquinolones) are as follows: diarrhea 5.0% vs. 6.2%; rash 3.5%  
762 vs. 1.1%; nausea 3.7% vs. 4.5%; headache 4.2% vs. 5.2%; abdominal pain 2.2% vs.  
763 2.2%; vomiting 1.6% vs. 2.0%; and dizziness 1.7% vs. 2.6%.

764 FACTIVE appears to have a low potential for photosensitivity. In clinical trials,  
765 treatment-related photosensitivity occurred in only 0.039% (3/7659) of patients.

766  
767 **Adverse Events with a Frequency of Less than 1%**

768 Additional drug-related adverse events (possibly or probably related) in the 8119 patients,  
769 with a frequency of >0.1% to ≤1% included: abdominal pain, anorexia, constipation,  
770 dermatitis, dizziness, dry mouth, dyspepsia, fatigue, flatulence, fungal infection, gastritis,  
771 genital moniliasis, genital pruritus, hyperglycemia, increased alkaline phosphatase,  
772 increased ALT, increased AST, increased creatine phosphokinase, insomnia, leukopenia,  
773 pruritus, somnolence, taste perversion, thrombocytopenia, urticaria, vaginitis, and  
774 vomiting.

775 Other adverse events reported from clinical trials which have potential clinical  
776 significance and which were considered to have a suspected relationship to the drug, that  
777 occurred in ≤0.1% of patients were: abnormal urine, abnormal vision, anemia, arthralgia,  
778 asthenia, back pain, bilirubinemia, dyspnea, eczema, eosinophilia, facial edema, flushing,  
779 gastroenteritis, granulocytopenia, hot flashes, increased GGT, increased non-protein  
780 nitrogen, leg cramps, moniliasis, myalgia, nervousness, non-specified gastrointestinal  
781 disorder, pain, pharyngitis, pneumonia, thrombocytopenia, tremor, vertigo.

782 In clinical trials of acute bacterial exacerbation of chronic bronchitis (ABECB)  
783 and community acquired pneumonia (CAP), the incidences of rash were as follows  
784 (Table 3):

785  
786 **Table 3. Incidence of Rash by Clinical Indication in Patients Treated with**  
787 **FACTIVE**  
788

	ABECB (5 days) N = 2284		CAP (5 days) N = 256		CAP (7 days) N = 643	
	n/N	%	n/N	%	n/N	%
Totals	27/2284	1.2	1/256	0.4	26/643	4.0
Females, < 40 years	NA*		1/37	2.7	8/88	9.1
Females, ≥ 40 years	16/1040	1.5	0/73	0	5/214	2.3
Males, < 40 years	NA*		0/65	0	5/101	5.0
Males, ≥ 40 years	11/1203	0.9	0/81	0	8/240	3.3

\* insufficient number of patients in this category for a meaningful analysis

789

790 (See **PRECAUTIONS**).

791

792 **Laboratory Changes:** The percentages of patients who received multiple doses of  
793 FACTIVE and had a laboratory abnormality are listed below. It is not known whether  
794 these abnormalities were related to FACTIVE or an underlying condition.

795 Clinical Chemistry: increased ALT (1.7%), increased AST (1.3%), increased  
796 creatine phosphokinase (0.7%), increased alkaline phosphatase (0.4%), increased total  
797 bilirubin (0.4%), increased potassium (0.3%), decreased sodium (0.2%), increased blood

---

798 urea nitrogen (0.3%), decreased albumin (0.3%), increased serum creatinine (0.2%),  
799 decreased calcium (0.1%), decreased total protein (0.1%), decreased potassium (0.1%),  
800 increased sodium (0.1%), increased lactate dehydrogenase (<0.1%) and increased  
801 calcium (<0.1%).

802 CPK elevations were noted infrequently: 0.7% in FACTIVE patients vs. 0.7% in  
803 the comparator patients.

804 Hematology: increased platelets (1.0%), decreased neutrophils (0.5%), increased  
805 neutrophils (0.5%), decreased hematocrit (0.3%), decreased hemoglobin (0.2%),  
806 decreased platelets (0.2%), decreased red blood cells (0.1%), increased hematocrit  
807 (0.1%), increased hemoglobin (0.1%), and increased red blood cells (0.1%).

808 In clinical studies, approximately 7% of the FACTIVE treated patients had  
809 elevated ALT values immediately prior to entry into the study. Of these patients,  
810 approximately 15% showed a further elevation of their ALT at the on-therapy visit and  
811 9% showed a further elevation at the end of therapy visit. None of these patients  
812 demonstrated evidence of hepatocellular jaundice. For the pooled comparators,  
813 approximately 6% of patients had elevated ALT values immediately prior to entry into  
814 the study. Of these patients, approximately 7% showed a further elevation of their ALT  
815 at the on-therapy visit and 4% showed a further elevation at the end of therapy visit.

816 In a clinical trial where 638 patients received either a single 640 mg dose of  
817 gemifloxacin or 250 mg BID of ciprofloxacin for 3 days, there was an increased  
818 incidence of ALT elevations in the gemifloxacin arm (3.9%) vs. the comparator arm  
819 (1.0%). In this study, two patients experienced ALT elevations of 8 to 10 times the upper  
820 limit of normal. These elevations were asymptomatic and reversible.

821

822 **Post-Marketing Adverse Reactions:** The majority of the post-marketing adverse events  
823 reported were cutaneous and most of these were rash. Some of these cutaneous adverse  
824 events were considered serious. The majority of rashes occurred in women and in  
825 patients under 40 years of age.

826

827 The following are additional adverse reactions reported during the post-marketing use of  
828 FACTIVE. Since these reactions are reported voluntarily from a population of uncertain  
829 size, it is impossible to reliably estimate their frequency or establish a causal relationship  
830 to FACTIVE exposure:

- 831 • anaphylactic reaction, erythema multiforme, skin exfoliation, facial swelling;
- 832 • hemorrhage, increased international normalized ratio (INR), retinal hemorrhage;
- 833 • peripheral edema;
- 834 • renal failure;
- 835 • prolonged QT, supraventricular tachycardia, syncope, transient ischemic attack;
- 836 • antibiotic-associated colitis.

837

---

838 **OVERDOSAGE**

839 Any signs or symptoms of overdosage should be treated symptomatically. No specific  
840 antidote is known. In the event of acute oral overdosage, the stomach should be emptied  
841 by inducing vomiting or by gastric lavage; the patient should be carefully observed and  
842 treated symptomatically with appropriate hydration maintained. Hemodialysis removes  
843 approximately 20 to 30% of an oral dose of gemifloxacin from plasma.

844 Mortality occurred at oral gemifloxacin doses of 1600 mg/kg in rats and 320  
845 mg/kg in mice. The minimum lethal intravenous doses in these species were 160 and 80  
846 mg/kg, respectively. Toxic signs after administration of a single high oral dose (400  
847 mg/kg) of gemifloxacin to rodents included ataxia, lethargy, piloerection, tremor, and  
848 clonic convulsions.

849

850 **DOSAGE AND ADMINISTRATION**

851 FACTIVE can be taken with or without food and should be swallowed whole with a  
852 liberal amount of liquid. The recommended dose of FACTIVE is 320 mg daily,  
853 according to the following table (Table 4).

854

855 **Table 4. Recommended Dosage Regimen of FACTIVE**

856 The clinical decision regarding the use of a 5 day or 7 day regimen should be guided by  
857 results of the initial sputum culture.

858

INDICATION	DOSE/DURATION
Acute bacterial exacerbation of chronic bronchitis	One 320 mg tablet daily for 5 days
Community-acquired pneumonia (of mild to moderate severity)	
<i>due to known or suspected S. pneumoniae, H. influenzae, M. pneumoniae, or C. pneumoniae infection</i>	One 320 mg tablet daily for 5 days
<i>due to known or suspected MDRSP*, K. pneumoniae, or M. catarrhalis infection</i>	One 320 mg tablet daily for 7 days

\*MDRSP, multi-drug resistant *Streptococcus pneumoniae*, includes isolates previously known as PRSP (penicillin-resistant *Streptococcus pneumoniae*), and are strains resistant to two or more of the following antibiotics: penicillin (MIC  $\geq 2$   $\mu\text{g/mL}$ ), 2nd generation cephalosporins (e.g., cefuroxime), macrolides, tetracyclines and trimethoprim/sulfamethoxazole.

859

860 The recommended dose and duration of FACTIVE should not be exceeded (see Table 2).

861

862 **Use in Renally Impaired Patients:** Dose adjustment in patients with creatinine clearance  
863 >40 mL/min is not required. Modification of the dosage is recommended for patients  
864 with creatinine clearance ≤40 mL/min. Table 5 provides dosage guidelines for use in  
865 patients with renal impairment:

866

867 **Table 5. Recommended Doses for Patients with Renal Impairment**

Creatinine Clearance (mL/min)	Dose
>40	See Usual Dosage
≤40	160 mg every 24 hours

868

869 Patients requiring routine hemodialysis or continuous ambulatory peritoneal dialysis  
870 (CAPD) should receive 160 mg every 24 hours.

871

872 When only the serum creatinine concentration is known, the following formula  
873 may be used to estimate creatinine clearance.

874

875 Men: Creatinine Clearance (mL/min) =  $\frac{\text{Weight (kg)} \times (140 - \text{age})}{72 \times \text{serum creatinine (mg/dL)}}$

876

877 Women: 0.85 x the value calculated for men

878

879 **Use in Hepatically Impaired Patients:** No dosage adjustment is recommended in  
880 patients with mild (Child-Pugh Class A), moderate (Child-Pugh Class B) or severe  
881 (Child-Pugh Class C) hepatic impairment.

882

883 **Use in Elderly:** No dosage adjustment is recommended.

884

## 885 HOW SUPPLIED

886 FACTIVE (gemifloxacin mesylate) is available as white to off-white, oval, film-coated  
887 tablets with breaklines and GE 320 debossed on both faces. Each tablet contains  
888 gemifloxacin mesylate equivalent to 320 mg of gemifloxacin.

889

890 320 mg Unit of Use (CR\*) 5's NDC 67707-320-05

891 320 mg Unit of Use (CR\*) 7's NDC 67707-320-07

892 \*Child Resistant

893

## 894 Storage

895 Store at 25°C (77°F); excursions permitted to 15°-30°C (59°-86°F) [see USP Controlled  
896 Room Temperature]. Protect from light.

897

## 898 ANIMAL PHARMACOLOGY

899 Quinolones have been shown to cause arthropathy in immature animals. Degeneration of  
900 articular cartilage occurred in juvenile dogs given at least 192 mg/kg/day gemifloxacin in  
901

902 a 28-day study (producing about 6 times the systemic exposure at the clinical dose), but  
903 not in mature dogs. There was no damage to the articular surfaces of joints in immature  
904 rats given repeated doses of up to 800 mg/kg/day.

905 Some quinolones have been reported to have proconvulsant properties that are  
906 potentiated by the concomitant administration of non-steroidal anti-inflammatory drugs  
907 (NSAIDs). Gemifloxacin alone had effects in tests of behavior or CNS interaction  
908 typically at doses of at least 160 mg/kg. No convulsions occurred in mice given the  
909 active metabolite of the NSAID, fenbufen, followed by 80 mg/kg gemifloxacin.

910 Dogs given 192 mg/kg/day (about 6 times the systemic exposure at the clinical  
911 dose) for 28 days, or 24 mg/kg/day (approximately equivalent to the systemic exposure at  
912 the clinical dose) for 13 weeks showed reversible increases in plasma ALT activities and  
913 local periportal liver changes associated with blockage of small bile ducts by crystals  
914 containing gemifloxacin.

915 Quinolones have been associated with prolongation of the electrocardiographic  
916 QT interval in dogs. Gemifloxacin produced no effect on the QT interval in dogs dosed  
917 orally to provide about 4 times human therapeutic plasma concentrations at C<sub>max</sub>, and  
918 transient prolongation after intravenous administration at more than 4 times human  
919 plasma levels at C<sub>max</sub>. Gemifloxacin exhibited weak activity in the cardiac I<sub>Kr</sub> (hERG)  
920 channel inhibition assay, having an IC<sub>50</sub> of approximately 270 μM.

921 Gemifloxacin, like many other quinolones, tends to crystallize at the alkaline pH  
922 of rodent urine, resulting in a nephropathy in rats that is reversible on drug withdrawal  
923 (oral no-effect dose 24 mg/kg/day).

924 Gemifloxacin was weakly phototoxic to hairless mice given a single 200 mg/kg  
925 oral dose and exposed to UVA radiation. However, no evidence of phototoxicity was  
926 observed at 100 mg/kg/day dosed orally for 13 weeks in a standard hairless mouse model,  
927 using simulated sunlight.

928

## 929 **CLINICAL STUDIES**

### 930 **Acute Bacterial Exacerbation of Chronic Bronchitis (ABECB)**

931 FACTIVE (320 mg once daily for 5 days) was evaluated for the treatment of acute  
932 bacterial exacerbation of chronic bronchitis in three pivotal double-blind, randomized,  
933 actively-controlled clinical trials (studies 068, 070, and 212). The primary efficacy  
934 parameter in these studies was the clinical response at follow-up (day 13 to 24). The  
935 results of the clinical response at follow-up for the principal ABECB studies demonstrate  
936 that FACTIVE 320 mg PO once daily for 5 days was at least as good as the comparators  
937 given for 7 days. The results are shown in Table 6 below.

938

939 **Table 6. Clinical Response at Follow-Up (Test of Cure): Pivotal ABECB Studies**

<b>Drug Regimen</b>	<b>Success Rate % (n/N)</b>	<b>Treatment Difference (95% CI)</b>
<b>Study 068</b>		
FACTIVE 320 mg x 5 days	86.0 (239/278)	1.2 (-4.7, 7.0)
Clarithromycin 500 mg BID x 7 days	84.8 (240/283)	
<b>Study 070</b>		
FACTIVE 320 mg x 5 days	93.6 (247/264)	0.4 (-3.9, 4.6)

Amoxicillin/clavulanate 500 mg/125 mg TID x 7 days	93.2 (248/266)	
<b>Study 212</b>		
FACTIVE 320 mg x 5 days	88.2 (134/152)	3.1 (-4.7, 10.7)
Levofloxacin 500 mg x 7 days	85.1 (126/148)	

940

941 **Community Acquired Pneumonia (CAP)**

942 **5 Day Treatment Regimen**

943 To evaluate the safety and efficacy of a 5-day course of FACTIVE, 510 outpatient and  
 944 hospitalized adults with clinically and radiologically determined mild to moderate  
 945 community-acquired pneumonia were clinically evaluated in a double-blind, randomized,  
 946 prospective, multicenter study comparing FACTIVE 320 mg for five days to  
 947 gemifloxacin 320 mg for seven days (Study OP-634-001).

948

949 Clinical success rates in the clinically evaluable population were 95.0% in the 5  
 950 day group and 92.1% in the 7 day group.

951

952 **Table 7. Clinical Response at Follow-Up (Test of Cure): Study OP-634-001**

Drug Regimen	Success Rate % (n/N)	Treatment Difference (95% CI)
<b>Study OP-634-001</b>		
FACTIVE 320 mg x 5 days	95.0 (230/242)	3.0 (-1.5, 7.4)
FACTIVE 320 mg x 7 days	92.1 (209/227)	

953

954 The microbiological efficacy of the 5-day regimen was documented for pathogens  
 955 listed in Table 8 below.

956

957 **Table 8. Bacterial Eradication by Pathogen for Patients Treated with FACTIVE in**  
 958 **Study OP-634-001**

959

Pathogen	5-day		7-day	
	n/N	%	n/N	%
<i>Streptococcus pneumoniae</i>	26/26	100	34/40	85.0
<i>Mycoplasma pneumoniae</i>	22/25	88.0	19/20	95.0
<i>Haemophilus influenzae</i>	21/22	95.5	18/18	100
<i>Chlamydia pneumoniae</i>	17/18	94.4	30/31	96.8

960

961 **7-Day Treatment Regimen**

962 Previous clinical studies evaluated the efficacy of FACTIVE in a 7-day treatment of CAP  
 963 in adults. This clinical program consisted of three double-blind, randomized, actively-  
 964 controlled clinical studies (studies 011, 012, and 049) and one open-label, actively-  
 965 controlled study (study 185). In addition, two uncontrolled studies (studies 061 and 287)  
 966 were conducted. Three of the studies, controlled study 011 and the uncontrolled studies,

967 had a fixed 7-day duration of treatment for FACTIVE. Controlled study 011 compared a  
 968 7-day course of FACTIVE with a 10-day treatment course of amoxicillin/clavulanate  
 969 (1g/125 mg TID) and clinical success rates were similar between treatment arms. The  
 970 results of comparative studies 049, 185, and 012 were supportive although treatment  
 971 duration could have been 7 to 14 days. The results of the clinical studies with a fixed  
 972 7-day duration of gemifloxacin are shown in Table 9:

973  
 974 **Table 9. Clinical Response at Follow-Up (Test of Cure): CAP Studies with a Fixed 7-**  
 975 **day Duration of Treatment**

Drug Regimen	Success Rate % (n/N)	Treatment Difference (95% CI)*
<b>Study 011</b>		
FACTIVE 320 mg x 7 days	88.7 (102/115)	1.1 (-7.3, 9.5)
Amoxicillin/clavulanate 1 g/125 mg TID x 10 days	87.6 (99/113)	
<b>Study 061</b>		
FACTIVE 320 mg x 7 days	91.7 (154/168)	(86.1, 95.2)
<b>Study 287</b>		
FACTIVE 320 mg x 7 days	89.8 (132/147)	(84.9, 94.7)

976 \*For uncontrolled studies, the 95% CI around the success rate is shown

977

978 The combined bacterial eradication rates for patients treated with a fixed 7-day  
 979 treatment regimen of FACTIVE are shown in Table 10:

980

981 **Table 10. Bacterial Eradication by Pathogen for Patients Treated with FACTIVE in**  
 982 **Studies with a Fixed 7-day Duration of Treatment**

Pathogen	n/N	%
<i>S. pneumoniae</i>	102/117	87.2
<i>M. pneumoniae</i>	40/42	95.2
<i>H. influenzae</i>	48/53	90.6
<i>C. pneumoniae</i>	43/45	95.6
<i>K. pneumoniae</i>	18/20	90.0
<i>M. catarrhalis</i>	11/12	91.7

983

984 **7-day Treatment Regimen of Community-Acquired Pneumonia Due to Multi-Drug**  
 985 **Resistant *Streptococcus pneumoniae* (MDRSP)**

986 FACTIVE was also effective in the treatment of CAP due to multi-drug resistant  
 987 *Streptococcus pneumoniae* (MDRSP\*). Of 35 patients with MDRSP treated for 7 days,  
 988 29 (82.9%) achieved clinical and bacteriological success at follow-up. The clinical and  
 989 bacteriological success for the 35 patients with MDRSP isolates are shown in Table 11.

990



991 \*MDRSP: multi-drug resistant *Streptococcus pneumoniae*, includes isolates  
 992 previously known as PRSP (penicillin-resistant *Streptococcus pneumoniae*), and are  
 993 strains resistant to two or more of the following antibiotics: penicillin (MIC  $\geq 2$   $\mu\text{g/mL}$ ),  
 994 2nd generation cephalosporins (e.g., cefuroxime), macrolides, tetracyclines and  
 995 trimethoprim/sulfamethoxazole.  
 996

997 **Table 11. Clinical and Bacteriological Success for 35 Patients Treated with**  
 998 **FACTIVE in CAP Studies with a 7-day Duration of Treatment for MDRSP**

Screening Susceptibility	Clinical Success		Bacteriological Success	
	n/N <sup>a</sup>	%	n/N <sup>b</sup>	%
Penicillin-resistant	15/16	93.8	15/16	93.8
2 <sup>nd</sup> generation cephalosporin-resistant	20/22	90.9	20/22	90.9
Macrolide-resistant <sup>c</sup>	24/28	85.7	23/28	82.1
Trimethoprim/sulfamethoxazole-resistant	23/26	88.5	23/26	88.5
Tetracycline-resistant	21/27	77.8	20/27	74.1

999

1000 <sup>a</sup>n = the number of patients successfully treated; N = number of patients with MDRSP

1001 <sup>b</sup>n = the number of bacteriological isolates successfully treated; N = number of isolates  
 1002 studied

1003 <sup>c</sup>Macrolide antibiotics tested include clarithromycin and erythromycin  
 1004

1005 Not all isolates were resistant to all antimicrobial classes tested. Success and  
 1006 eradication rates are summarized in the Table 10 below.  
 1007

1008 **Table 12. Resistant *Streptococcus pneumoniae* Clinical Success and Bacteriological**  
 1009 **Eradication Rates**  
 1010

<i>S. pneumoniae</i> with MDRSP	Clinical Cure Rate		Bacteriological Eradication Rate	
	n/N	%	n/N	%
Resistant to 2 antimicrobials	8/11	72.7	7/11	63.6
Resistant to 3 antimicrobials	5/7	71.4	5/7	71.4
Resistant to 4 antimicrobials	8/9	88.9	8/9	88.9
Resistant to 5 antimicrobials	8/8	100	8/8	100
Bacteremia with MDRSP	3/3	100	3/3	100

1011

1012

1013

1014

---

## 1015 **Clinical Safety Study of Rash**

1016  
1017 To further characterize gemifloxacin-associated rash, which in early clinical  
1018 studies appeared to be associated with age less than 40 and female gender, a clinical  
1019 pharmacology study was conducted. The study enrolled 1,011 healthy female volunteers  
1020 less than 40 years of age. Subjects were randomized in a 5:1 ratio to receive either  
1021 FACTIVE 320 mg PO daily (819 subjects) or ciprofloxacin 500 mg PO twice daily for 10  
1022 days (164 subjects). This study was designed to enroll subjects at high risk for rash  
1023 (women <40 years of age and dosing beyond the recommended duration of therapy for  
1024 FACTIVE [10 days]), and over estimates the risk to patients taking FACTIVE as  
1025 prescribed. Subjects who received FACTIVE were 7 times more likely to develop rash  
1026 than those who received ciprofloxacin. Of the 260 rashes in subjects receiving  
1027 FACTIVE, the majority of the rashes were maculopapular and of mild to moderate  
1028 severity; 7% of the rashes were reported as severe, and severity appeared to correlate  
1029 with the extent of the rash. In 68% of the subjects reporting a severe rash and  
1030 approximately 25% of all those reporting rash, >60% of the body surface area was  
1031 involved; the characteristics of the rash were otherwise indistinguishable from those  
1032 subjects reporting a mild rash. The histopathology was consistent with the clinical  
1033 observation of uncomplicated exanthematous morbilliform eruption. Approximately  
1034 11% of the rashes were described as being “urticaria-like”. There were no documented  
1035 cases of hypersensitivity syndrome or findings suggestive of angioedema or other serious  
1036 cutaneous reactions.

1037 The majority of rashes (81.9%) occurred on days 8 through 10 of the planned 10  
1038 day course of FACTIVE; 2.7% of rash events occurred within one day of the start of  
1039 dosing. The median duration of rash was 6 days. The rash resolved without treatment in  
1040 the majority of subjects. Approximately 19% received antihistamines and 5% received  
1041 steroids, although the therapeutic benefit of these therapies is uncertain.

1042 In the second part of this study after a 4 to 6 week wash out period, subjects  
1043 developing a rash on FACTIVE were treated with ciprofloxacin (n=136) or placebo  
1044 (n=50); 5.9% developed rash when treated with ciprofloxacin and 2.0% developed rash  
1045 when treated with placebo. The cross sensitization rate to other fluoroquinolones was not  
1046 evaluated in this clinical study. There was no evidence of sub-clinical sensitization to  
1047 FACTIVE on a second exposure (i.e., subjects who had not developed a rash to  
1048 FACTIVE in the first part of the study were not at higher risk of developing a rash to  
1049 FACTIVE with a second exposure).

1050 There was no relationship between the incidence of rash and systemic exposure  
1051 (C<sub>max</sub> and AUC) to either gemifloxacin or its major metabolite, N-acetyl gemifloxacin.

1052  
1053 **REFERENCES:** 1. Clinical and Laboratory Standards Institute. Methods for Dilution  
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1060

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1061 DATE OF REVISION May 2007  
1062 © Oscient Pharmaceuticals Corporation 2007  
1063 FACTIVE is a registered trademark of LG Life Sciences.

1064 **Rx only**

1065  
1066 Manufactured for:  
1067 Oscient Pharmaceuticals  
1068 Waltham, MA 02451-1478 USA

1069  
1070 Licensed from LG Life Sciences, Ltd. Seoul, Korea

1071  
1072

1073 **Patient Information**

1074

1075 **FACTIVE®**  
1076 **(gemifloxacin mesylate) Tablets**

1077

1078 This leaflet summarizes the most important information about FACTIVE. Read the  
1079 Patient Information that comes with FACTIVE each time you get a new prescription.  
1080 There may be new information. This leaflet does not list all benefits and risks of  
1081 treatment and does not take the place of talking with your healthcare provider about your  
1082 condition or your treatment.

1083

1084 **What is FACTIVE?**

1085 FACTIVE is an antibiotic. It is used to treat adults 18 years or older with bronchitis or  
1086 pneumonia (lung infections) caused by certain bacteria (germs).

1087

1088 Sometimes, other germs called viruses infect the lungs. The common cold is a virus.  
1089 FACTIVE, like other antibiotics, does not treat viruses.

1090

1091 **Who should not take FACTIVE?**

- 1092 • **Do not take FACTIVE if you are allergic to any of the ingredients in FACTIVE**  
1093 **or to any antibiotic called a “quinolone”.** If you develop hives, difficulty  
1094 breathing, or other symptoms of a severe allergic reaction, seek emergency treatment  
1095 right away. If you develop a skin rash, stop taking FACTIVE and call your  
1096 healthcare professional. The ingredients in FACTIVE are listed at the end of this  
1097 leaflet. Ask your healthcare provider or pharmacist if you need a list of quinolone  
1098 antibiotics.

1099

1100 **FACTIVE may not be right for you. Tell your healthcare provider if you:**

- 1101 • are pregnant, planning to become pregnant, or are breast feeding. The effects of  
1102 FACTIVE on unborn children and nursing infants are unknown;
- 1103 • or any family members have a rare heart condition known as congenital prolongation  
1104 of the QTc interval;
- 1105 • have low potassium or magnesium levels;
- 1106 • have a slow heart beat called bradycardia;

- 
- 1107 • have had a recent heart attack;  
1108 • have a history of seizures or epilepsy;  
1109 • have kidney problems.

1110  
1111 FACTIVE has not been studied in children under the age of 18. Quinolones, such as  
1112 FACTIVE may cause joint problems (arthropathy) in children.

1113  
1114 **What about other medicines I am taking?**

1115 Tell your healthcare provider about all the medicines you take including prescription and  
1116 nonprescription medicines, vitamins, and dietary supplements. FACTIVE and other  
1117 medicines may affect each other, causing serious side effects. **Be sure to tell your**  
1118 **healthcare provider if you take:**

- 1119
- 1120 • medicines for your heart rhythm called “antiarrhythmics”
  - 1121 • erythromycin
  - 1122 • medicines for your mental health called “antipsychotics” or “tricyclic  
1123 antidepressants”
  - 1124 • medicines called “corticosteroids”, taken by mouth or by injection
  - 1125 • medicines called “water pills” (diuretics) such as furosemide and  
1126 hydrochlorothiazide;
  - 1127 • medicines to thin your blood (called oral anticoagulants) such as Coumadin<sup>®</sup> or  
1128 warfarin.

1129  
1130 **How should I take FACTIVE?**

- 1131 • Take 1 FACTIVE tablet a day for 5 or 7 days, exactly as prescribed.
- 1132 • Take FACTIVE at the same time each day.
- 1133 • FACTIVE can be taken with or without food.
- 1134 • Swallow the FACTIVE tablet whole, and drink plenty of fluids with it. Do not chew  
1135 the FACTIVE tablet.
- 1136 • If you miss a dose of FACTIVE, take it as soon as you remember. **Do not take more**  
1137 **than 1 dose of FACTIVE in a day.**
- 1138 • To make sure all bacteria are killed, take all the medicine that was prescribed for you  
1139 even if you begin to feel better.
- 1140 • Call your healthcare provider if your condition does not improve while taking  
1141 FACTIVE.

1142  
1143 **Do not take the following medicines within 3 hours before FACTIVE or 2 hours**  
1144 **after FACTIVE.** They may interfere with the absorption of FACTIVE and may prevent  
1145 it from working properly:

- 1146
- 1147 • antacids that contain magnesium or aluminum
  - 1148 • ferrous sulfate (iron)
  - 1149 • multivitamin that contains zinc or other metals
  - 1150 • Videx<sup>®</sup> (didanosine)

1151

---

1152 **FACTIVE should be taken at least 2 hours before sucralfate.**

1153

1154 **What are possible side effects of FACTIVE?**

1155 FACTIVE is generally well tolerated. The most common side effects with FACTIVE  
1156 include diarrhea, rash, nausea, headache, vomiting, stomach pain, dizziness, and a change  
1157 in the way things taste in your mouth.

1158

1159 Rash occurs more commonly in women, especially those on hormone replacement  
1160 therapy and anyone under 40. The likelihood of getting a rash increases if FACTIVE is  
1161 taken for longer than 7 days. The rash is usually mild to moderate, but may occasionally  
1162 be severe. If you get a rash while taking FACTIVE, stop FACTIVE, and call your  
1163 healthcare provider right away.

1164

1165 **FACTIVE and other quinolone antibiotics may cause the following serious side**  
1166 **effects:**

1167

1168 • **a rare heart problem known as prolongation of the QTc interval.** This condition  
1169 can cause an abnormal heartbeat and result in sudden death. The chances of this  
1170 event are increased in those with a family history of prolonged QT interval, low  
1171 potassium (hypokalemia), and who are taking drugs to control heart rhythm, called  
1172 Class IA (quinidine, procainamide) or Class III (amiodarone, sotalol) antiarrhythmic  
1173 agents.

1174

1175 You should call your healthcare provider right away if you have any symptoms of  
1176 prolongation of the QTc interval including:

- 1177 • heart palpitations (a change in the way your heart beats)
- 1178 • a loss of consciousness (fainting spells)

1179

1180 • **allergic reactions.** Get medical help right away if you develop hives, trouble  
1181 breathing, wheezing, or other symptoms of a severe allergic reaction.

1182

1183 • **tendon problems including pain, swelling (tendonitis) or rupture (“tears”) of**  
1184 **Achilles, shoulder or hand tendons.** The risk for tendon problems is higher if you  
1185 are over 65 years old and/or are taking corticosteroids. If you experience pain,  
1186 swelling, or rupture of a tendon, you should stop taking FACTIVE, avoid exercise  
1187 and strenuous use of the affected area, and call your healthcare provider;

1188

1189 • **diarrhea** that usually ends after treatment is a common problem caused by  
1190 antibiotics. A more serious form of diarrhea with inflammation of the colon  
1191 (pseudomembranous colitis) can occur during or up to 2 months after the use of  
1192 antibiotics. This has been reported with all antibiotics including with FACTIVE. If  
1193 you develop a watery and bloody stool with or without stomach cramps and fever,  
1194 contact your physician as soon as possible.

1195

- 
- 1196 • **central nervous system problems** including body shakes (tremors), restless feeling,  
1197 lightheaded feelings, confusion, and hallucinations (seeing or hearing things that are  
1198 not there);  
1199
- 1200 • **dizziness.** FACTIVE can make you dizzy. Do not drive or operate heavy machinery  
1201 until you know how FACTIVE affects you.  
1202
- 1203 • **phototoxicity.** FACTIVE may rarely make your skin sunburn more easily. Do not  
1204 use a sunlamp or tanning bed while taking FACTIVE. Use a sunscreen and wear  
1205 protective clothing if you must be out in the sun.  
1206

1207 These are not all the side effects you may experience with FACTIVE. If you get any side  
1208 effects that concern you, call your healthcare provider.  
1209

#### 1210 **How should I store Factive?**

- 1211 • Store FACTIVE at room temperature between 59° and 86° F (15° to 30° C). Protect  
1212 from light.  
1213 • **Keep FACTIVE and all medicines out of the reach of children.**  
1214

#### 1215 **General information about the safe and effective use of FACTIVE:**

1216  
1217 Medicines are sometimes prescribed for conditions other than those described in patient  
1218 information leaflets. Do not use FACTIVE for a condition for which it was not  
1219 prescribed. Do not give FACTIVE to other people, even if they have the same symptoms  
1220 that you have. It may harm them.  
1221

#### 1222 **What are the ingredients in FACTIVE?**

1223 Active ingredient: gemifloxacin  
1224 Inactive ingredients: crospovidone, hydroxypropyl methycellulose, magnesium stearate,  
1225 microcrystalline cellulose, polyethylene glycol, povidone, titanium dioxide.  
1226

1227 FACTIVE tablets are white to off-white and imprinted with GE 320 on both sides.  
1228

1229 This leaflet summarizes the most important information about FACTIVE. If you would  
1230 like more information, talk with your healthcare provider. You can ask your healthcare  
1231 provider or pharmacist for information about FACTIVE that is written for healthcare  
1232 professionals. For more information, visit our website at [www.factive.com](http://www.factive.com).  
1233

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1239 Manufactured for:

1240 Oscient Pharmaceuticals

1241 Waltham, MA 02451-1478 USA

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