1	< Roche >
2	LARIAM [®]
3	brand of
4	mefloquine hydrochloride
5	TABLETS
6	R _X only
7 8 9 10	DESCRIPTION Lariam (mefloquine hydrochloride) is an antimalarial agent available as 250-mg tablets of mefloquine hydrochloride (equivalent to 228.0 mg of the free base) for oral administration.
11 12 13 14 15	Mefloquine hydrochloride is a 4-quinolinemethanol derivative with the specific chemical name of (R^*, S^*) - (\pm) - α -2-piperidinyl-2,8-bis (trifluoromethyl)-4-quinolinemethanol hydrochloride. It is a 2-aryl substituted chemical structural analog of quinine. The drug is a white to almost white crystalline compound, slightly soluble in water.
16 17	Mefloquine hydrochloride has a calculated molecular weight of 414.78 and the following structural formula:
	CF ₃ N CF ₃ HC—OH HC—NH
18	
19 20 21	The inactive ingredients are ammonium-calcium alginate, corn starch, crospovidone, lactose, magnesium stearate, microcrystalline cellulose, poloxamer #331, and talc.
22	CLINICAL PHARMACOLOGY
23	Pharmacokinetics
24 25 26 27 28 29	Absorption The absolute oral bioavailability of mefloquine has not been determined since an intravenous formulation is not available. The bioavailability of the tablet formation compared with an oral solution was over 85%. The presence of food significantly enhances the rate and extent of absorption, leading to about a 40% increase in bioavailability. In healthy volunteers, plasma concentrations

- 30 peak 6 to 24 hours (median, about 17 hours) after a single dose of Lariam. In a
- 31 similar group of volunteers, maximum plasma concentrations in μg/L are
- 32 roughly equivalent to the dose in milligrams (for example, a single 1000 mg
- dose produces a maximum concentration of about 1000 µg/L). In healthy
- volunteers, a dose of 250 mg once weekly produces maximum steady-state
- 35 plasma concentrations of 1000 to 2000 μg/L, which are reached after 7 to 10
- 36 weeks.
- 37 Distribution
- 38 In healthy adults, the apparent volume of distribution is approximately 20
- 39 L/kg, indicating extensive tissue distribution. Mefloquine may accumulate in
- 40 parasitized erythrocytes. Experiments conducted in vitro with human blood
- 41 using concentrations between 50 and 1000 mg/mL showed a relatively
- 42 constant erythrocyte-to-plasma concentration ratio of about 2 to 1. The
- 43 equilibrium reached in less than 30 minutes was found to be reversible.
- 44 Protein binding is about 98%.
- 45 Mefloquine crosses the placenta. Excretion into breast milk appears to be
- 46 minimal (see **PRECAUTIONS: Nursing Mothers**).
- 47 Metabolism
- 48 Two metabolites have been identified in humans. The main metabolite, 2,8-
- 49 bis-trifluoromethyl-4-quinoline carboxylic acid, is inactive in Plasmodium
- 50 falciparum. In a study in healthy volunteers, the carboxylic acid metabolite
- 51 appeared in plasma 2 to 4 hours after a single oral dose. Maximum plasma
- 52 concentrations, which were about 50% higher than those of mefloquine, were
- reached after 2 weeks. Thereafter, plasma levels of the main metabolite and
- 54 mefloquine declined at a similar rate. The area under the plasma
- 55 concentration-time curve (AUC) of the main metabolite was 3 to 5 times
- larger than that of the parent drug. The other metabolite, an alcohol, was
- 57 present in minute quantities only.
- 58 Elimination
- 59 In several studies in healthy adults, the mean elimination half-life of
- 60 mefloquine varied between 2 and 4 weeks, with an average of about 3 weeks.
- Total clearance, which is essentially hepatic, is in the order of 30 mL/min.
- There is evidence that mefloquine is excreted mainly in the bile and feces. In
- volunteers, urinary excretion of unchanged mefloquine and its main
- 64 metabolite under steady-state condition accounted for about 9% and 4% of the
- dose, respectively. Concentrations of other metabolites could not be measured
- in the urine.

- 67 Pharmacokinetics in Special Clinical Situations
- 68 *Children and the Elderly*
- No relevant age-related changes have been observed in the pharmacokinetics
- of mefloquine. Therefore, the dosage for children has been extrapolated from
- 71 the recommended adult dose.
- No pharmacokinetic studies have been performed in patients with renal
- 73 insufficiency since only a small proportion of the drug is eliminated renally.
- 74 Mefloquine and its main metabolite are not appreciably removed by
- 75 hemodialysis. No special chemoprophylactic dosage adjustments are indicated
- 76 for dialysis patients to achieve concentrations in plasma similar to those in
- healthy persons.
- Although clearance of mefloquine may increase in late pregnancy, in general,
- 79 pregnancy has no clinically relevant effect on the pharmacokinetics of
- 80 mefloquine.
- 81 The pharmacokinetics of mefloquine may be altered in acute malaria.
- 82 Pharmacokinetic differences have been observed between various ethnic
- 83 populations. In practice, however, these are of minor importance compared
- with host immune status and sensitivity of the parasite.
- 85 During long-term prophylaxis (>2 years), the trough concentrations and the
- 86 elimination half-life of mefloquine were similar to those obtained in the same
- 87 population after 6 months of drug use, which is when they reached steady
- 88 state.
- 89 In vitro and in vivo studies showed no hemolysis associated with glucose-6-
- 90 phosphate dehydrogenase deficiency (see **ANIMAL TOXICOLOGY**).

91 **Microbiology**

- 92 Mechanism of Action
- 93 Mefloquine is an antimalarial agent which acts as a blood schizonticide. Its
- 94 exact mechanism of action is not known.
- 95 Activity In Vitro and In Vivo
- 96 Mefloquine is active against the erythrocytic stages of *Plasmodium* species
- 97 (see **INDICATIONS AND USAGE**). However, the drug has no effect against
- 98 the exoerythrocytic (hepatic) stages of the parasite. Mefloquine is effective
- 99 against malaria parasites resistant to chloroquine (see **INDICATIONS AND**
- 100 **USAGE**).
- 101 Drug Resistance
- Strains of *P. falciparum* with decreased susceptibility to mefloquine can be
- selected in vitro or in vivo. Resistance of *P. falciparum* to mefloquine has

- been reported in areas of multi-drug resistance in South East Asia. Increased
- incidences of resistance have also been reported in other parts of the world.
- 106 Cross-Resistance
- 107 Cross-resistance between mefloquine and halofantrine and cross-resistance
- between mefloquine and quinine have been observed in some regions.

109 INDICATIONS AND USAGE

110 Treatment of Acute Malaria Infections

- 111 Lariam is indicated for the treatment of mild to moderate acute malaria caused
- by mefloquine-susceptible strains of P. falciparum (both chloroquine-
- susceptible and resistant strains) or by *Plasmodium vivax*. There are
- insufficient clinical data to document the effect of mefloquine in malaria
- caused by *P. ovale* or *P. malariae*.
- Note: Patients with acute P. vivax malaria, treated with Lariam, are at
- high risk of relapse because Lariam does not eliminate exoerythrocytic
- (hepatic phase) parasites. To avoid relapse, after initial treatment of the
- acute infection with Lariam, patients should subsequently be treated
- with an 8-aminoquinoline derivative (eg, primaquine).

Prevention of Malaria

- Lariam is indicated for the prophylaxis of *P. falciparum* and *P. vivax* malaria
- infections, including prophylaxis of chloroquine-resistant strains of P.
- 124 falciparum.

121

125 **CONTRAINDICATIONS**

- 126 Use of Lariam is contraindicated in patients with a known hypersensitivity to
- mefloquine or related compounds (eg, quinine and quinidine) or to any of the
- excipients contained in the formulation. Lariam should not be prescribed for
- prophylaxis in patients with active depression, a recent history of depression,
- generalized anxiety disorder, psychosis, or schizophrenia or other major
- psychiatric disorders, or with a history of convulsions.

132 WARNINGS

- 133 In case of life-threatening, serious or overwhelming malaria infections
- due to *P. falciparum*, patients should be treated with an intravenous
- antimalarial drug. Following completion of intravenous treatment,
- 136 Lariam may be given to complete the course of therapy.
- Data on the use of halofantrine subsequent to administration of Lariam
- suggest a significant, potentially fatal prolongation of the OTc interval of
- the ECG. Therefore, halofantrine must not be given simultaneously with
- or subsequent to Lariam. No data are available on the use of Lariam after
- 141 halofantrine (see PRECAUTIONS: Drug Interactions).

- 142 Mefloquine may cause psychiatric symptoms in a number of patients,
- ranging from anxiety, paranoia, and depression to hallucinations and
- psychotic behavior. On occasions, these symptoms have been reported to
- continue long after mefloquine has been stopped. Rare cases of suicidal
- ideation and suicide have been reported though no relationship to drug
- administration has been confirmed. To minimize the chances of these
- 148 adverse events, mefloquine should not be taken for prophylaxis in
- 149 patients with active depression or with a recent history of depression,
- generalized anxiety disorder, psychosis, or schizophrenia or other major
- psychiatric disorders. Lariam should be used with caution in patients
- with a previous history of depression.
- During prophylactic use, if psychiatric symptoms such as acute anxiety,
- depression, restlessness or confusion occur, these may be considered
- prodromal to a more serious event. In these cases, the drug must be
- discontinued and an alternative medication should be substituted.
- 157 Concomitant administration of Lariam and quinine or quinidine may
- 158 produce electrocardiographic abnormalities.
- 159 Concomitant administration of Lariam and quinine or chloroquine may
- increase the risk of convulsions.

161 **PRECAUTIONS**

- 162 General
- Hypersensitivity reactions ranging from mild cutaneous events to anaphylaxis
- 164 cannot be predicted.
- In patients with epilepsy, Lariam may increase the risk of convulsions. The
- drug should therefore be prescribed only for curative treatment in such
- patients and only if there are compelling medical reasons for its use (see
- 168 **PRECAUTIONS: Drug Interactions**).
- 169 Caution should be exercised with regard to activities requiring alertness and
- 170 fine motor coordination such as driving, piloting aircraft, operating
- machinery, and deep-sea diving, as dizziness, a loss of balance, or other
- disorders of the central or peripheral nervous system have been reported
- during and following the use of Lariam. These effects may occur after therapy
- is discontinued due to the long half-life of the drug. Lariam should be used
- with caution in patients with psychiatric disturbances because mefloquine use
- 176 has been associated with emotional disturbances (see ADVERSE
- 177 **REACTIONS**).
- 178 In patients with impaired liver function the elimination of mefloquine may be
- prolonged, leading to higher plasma levels.
- 180 This drug has been administered for longer than 1 year. If the drug is to be
- administered for a prolonged period, periodic evaluations including liver

- 182 function tests should be performed. Although retinal abnormalities seen in
- humans with long-term chloroquine use have not been observed with 183
- 184 mefloquine use, long-term feeding of mefloquine to rats resulted in dose-
- 185 related ocular lesions (retinal degeneration, retinal edema and lenticular
- 186 opacity at 12.5 mg/kg/day and higher) (see ANIMAL TOXICOLOGY).
- 187 Therefore, periodic ophthalmic examinations are recommended.
- 188 Parenteral studies in animals show that mefloquine, a myocardial depressant,
- 189 possesses 20% of the antifibrillatory action of quinidine and produces 50% of
- 190 the increase in the PR interval reported with quinine. The effect of mefloquine
- 191 on the compromised cardiovascular system has not been evaluated. However,
- 192 transitory and clinically silent ECG alterations have been reported during the
- 193 use of mefloquine. Alterations included sinus bradycardia, sinus arrhythmia,
- 194 first degree AV-block, prolongation of the QTc interval and abnormal T
- 195 waves (see also cardiovascular effects under PRECAUTIONS: Drug
- 196 Interactions and ADVERSE REACTIONS). The benefits of Lariam therapy
- 197 should be weighed against the possibility of adverse effects in patients with
- 198 cardiac disease.

199 **Laboratory Tests**

- 200 Periodic evaluation of hepatic function should be performed during prolonged
- 201 prophylaxis.

202 **Information for Patients**

- 203 Medication Guide: As required by law, a Lariam Medication Guide is
- 204 supplied to patients when Lariam is dispensed. An information wallet card is
- 205 also supplied to patients when Lariam is dispensed. Patients should be
- 206 instructed to read the Medication Guide when Lariam is received and to carry
- 207 the information wallet card with them when they are taking Lariam. The
- 208 complete texts of the Medication Guide and information wallet card are
- 209 reprinted at the end of this document.
- 210 Patients should be advised:
- 211 that malaria can be a life-threatening infection in the traveler;
- 212 that Lariam is being prescribed to help prevent or treat this serious 213 infection;
- 214 that in a small percentage of cases, patients are unable to take this
- 215 medication because of side effects, and it may be necessary to change
- 216 medications;
- 217 that when used as prophylaxis, the first dose of Lariam should be taken 1 218 week prior to arrival in an endemic area;
- 219 that if the patients experience psychiatric symptoms such as acute anxiety,
- 220 depression, restlessness or confusion, these may be considered prodromal
- 221 to a more serious event. In these cases, the drug must be discontinued and
- 222 an alternative medication should be substituted;

- that no chemoprophylactic regimen is 100% effective, and protective clothing, insect repellents, and bednets are important components of
- 225 malaria prophylaxis;
- to seek medical attention for any febrile illness that occurs after return
 from a malarious area and to inform their physician that they may have
 been exposed to malaria.

229 **Drug Interactions**

- 230 Drug-drug interactions with Lariam have not been explored in detail. There is
- one report of cardiopulmonary arrest, with full recovery, in a patient who was
- taking a beta blocker (propranolol) (see **PRECAUTIONS: General**). The
- 233 effects of mefloquine on the compromised cardiovascular system have not
- been evaluated. The benefits of Lariam therapy should be weighed against the
- possibility of adverse effects in patients with cardiac disease.
- Because of the danger of a potentially fatal prolongation of the OTc interval.
- halofantrine must not be given simultaneously with or subsequent to Lariam
- (see **WARNINGS**).
- 239 Concomitant administration of Lariam and other related compounds (eg,
- 240 quinine, quinidine and chloroquine) may produce electrocardiographic
- abnormalities and increase the risk of convulsions (see WARNINGS). If
- these drugs are to be used in the initial treatment of severe malaria, Lariam
- 243 administration should be delayed at least 12 hours after the last dose. There is
- evidence that the use of halofantrine after mefloquine causes a significant
- lengthening of the QTc interval. Clinically significant QTc prolongation has
- 246 not been found with mefloquine alone.
- 247 This appears to be the only clinically relevant interaction of this kind with
- Lariam, although theoretically, coadministration of other drugs known to alter
- 249 cardiac conduction (eg, anti-arrhythmic or beta-adrenergic blocking agents,
- 250 calcium channel blockers, antihistamines or H₁-blocking agents, tricyclic
- antidepressants and phenothiazines) might also contribute to a prolongation of
- 252 the QTc interval. There are no data that conclusively establish whether the
- 253 concomitant administration of mefloquine and the above listed agents has an
- 254 effect on cardiac function.
- 255 In patients taking an anticonvulsant (eg, valproic acid, carbamazepine,
- 256 phenobarbital or phenytoin), the concomitant use of Lariam may reduce
- seizure control by lowering the plasma levels of the anticonvulsant. Therefore,
- 258 patients concurrently taking antiseizure medication and Lariam should have
- 259 the blood level of their antiseizure medication monitored and the dosage
- adjusted appropriately (see **PRECAUTIONS: General**).
- 261 When Lariam is taken concurrently with oral live typhoid vaccines,
- attenuation of immunization cannot be excluded. Vaccinations with attenuated
- 263 live bacteria should therefore be completed at least 3 days before the first dose
- of Lariam.

- No other drug interactions are known. Nevertheless, the effects of Lariam on
- 266 travelers receiving comedication, particularly diabetics or patients using
- anticoagulants, should be checked before departure.
- 268 In clinical trials, the concomitant administration of sulfadoxine and
- 269 pyrimethamine did not alter the adverse reaction profile.

270 Carcinogenesis, Mutagenesis, Impairment of Fertility

- 271 Carcinogenesis
- 272 The carcinogenic potential of mefloquine was studied in rats and mice in 2-
- year feeding studies at doses of up to 30 mg/kg/day. No treatment-related
- increases in tumors of any type were noted.
- 275 Mutagenesis
- 276 The mutagenic potential of mefloquine was studied in a variety of assay
- 277 systems including: Ames test, a host-mediated assay in mice, fluctuation tests
- and a mouse micronucleus assay. Several of these assays were performed with
- and without prior metabolic activation. In no instance was evidence obtained
- 280 for the mutagenicity of mefloquine.
- 281 Impairment of Fertility
- Fertility studies in rats at doses of 5, 20, and 50 mg/kg/day of mefloquine have
- demonstrated adverse effects on fertility in the male at the high dose of 50
- 284 mg/kg/day, and in the female at doses of 20 and 50 mg/kg/day.
- 285 Histopathological lesions were noted in the epididymides from male rats at
- doses of 20 and 50 mg/kg/day. Administration of 250 mg/week of mefloquine
- 287 (base) in adult males for 22 weeks failed to reveal any deleterious effects on
- 288 human spermatozoa.

Pregnancy

289

- 290 Teratogenic Effects
- 291 Pregnancy Category C. Mefloquine has been demonstrated to be teratogenic
- in rats and mice at a dose of 100 mg/kg/day. In rabbits, a high dose of 160
- 293 mg/kg/day was embryotoxic and teratogenic, and a dose of 80 mg/kg/day was
- 294 teratogenic but not embryotoxic. There are no adequate and well-controlled
- studies in pregnant women. However, clinical experience with Lariam has not
- 296 revealed an embryotoxic or teratogenic effect. Mefloquine should be used
- during pregnancy only if the potential benefit justifies the potential risk to the
- 298 fetus. Women of childbearing potential who are traveling to areas where
- malaria is endemic should be warned against becoming pregnant. Women of
- 300 childbearing potential should also be advised to practice contraception during
- 301 malaria prophylaxis with Lariam and for up to 3 months thereafter. However,
- in the case of unplanned pregnancy, malaria chemoprophylaxis with Lariam is
- 303 not considered an indication for pregnancy termination.

304 **Nursing Mothers**

- 305 Mefloquine is excreted in human milk in small amounts, the activity of which
- is unknown. Based on a study in a few subjects, low concentrations (3% to
- 307 4%) of mefloquine were excreted in human milk following a dose equivalent
- 308 to 250 mg of the free base. Because of the potential for serious adverse
- 309 reactions in nursing infants from mefloquine, a decision should be made
- 310 whether to discontinue the drug, taking into account the importance of the
- 311 drug to the mother.

312 **Pediatric Use**

- 313 Use of Lariam to treat acute, uncomplicated *P. falciparum* malaria in pediatric
- 314 patients is supported by evidence from adequate and well-controlled studies of
- 315 Lariam in adults with additional data from published open-label and
- 316 comparative trials using Lariam to treat malaria caused by *P. falciparum* in
- 317 patients younger than 16 years of age. The safety and effectiveness of Lariam
- 318 for the treatment of malaria in pediatric patients below the age of 6 months
- 319 have not been established.
- 320 In several studies, the administration of Lariam for the treatment of malaria
- was associated with early vomiting in pediatric patients. Early vomiting was
- 322 cited in some reports as a possible cause of treatment failure. If a second dose
- 323 is not tolerated, the patient should be monitored closely and alternative
- 324 malaria treatment considered if improvement is not observed within a
- reasonable period of time (see **DOSAGE AND ADMINISTRATION**).

326 Geriatric Use

- 327 Clinical studies of Lariam did not include sufficient numbers of subjects aged
- 328 65 and over to determine whether they respond differently from younger
- 329 subjects. Other reported clinical experience has not identified differences in
- 330 responses between the elderly and younger patients. Since
- 331 electrocardiographic abnormalities have been observed in individuals treated
- with Lariam (see **PRECAUTIONS**) and underlying cardiac disease is more
- prevalent in elderly than in younger patients, the benefits of Lariam therapy
- should be weighed against the possibility of adverse cardiac effects in elderly
- 335 patients.

ADVERSE REACTIONS

337 Clinical

336

- 338 At the doses used for treatment of acute malaria infections, the symptoms
- 339 possibly attributable to drug administration cannot be distinguished from
- those symptoms usually attributable to the disease itself.
- 341 Among subjects who received mefloquine for prophylaxis of malaria, the
- most frequently observed adverse experience was vomiting (3%). Dizziness,

- 343 syncope, extrasystoles and other complaints affecting less than 1% were also
- 344 reported.
- 345 Among subjects who received mefloquine for treatment, the most frequently
- 346 observed adverse experiences included: dizziness, myalgia, nausea, fever,
- headache, vomiting, chills, diarrhea, skin rash, abdominal pain, fatigue, loss of
- 348 appetite, and tinnitus. Those side effects occurring in less than 1% included
- 349 bradycardia, hair loss, emotional problems, pruritus, asthenia, transient
- 350 emotional disturbances and telogen effluvium (loss of resting hair). Seizures
- 351 have also been reported.
- 352 Two serious adverse reactions were cardiopulmonary arrest in one patient
- 353 shortly after ingesting a single prophylactic dose of mefloquine while
- concomitantly using propranolol (see **PRECAUTIONS: Drug Interactions**),
- and encephalopathy of unknown etiology during prophylactic mefloquine
- 356 administration. The relationship of encephalopathy to drug administration
- 357 could not be clearly established.

358 **Postmarketing**

- 359 Postmarketing surveillance indicates that the same kind of adverse
- experiences are reported during prophylaxis, as well as acute treatment.
- 361 The most frequently reported adverse events are nausea, vomiting, loose
- stools or diarrhea, abdominal pain, dizziness or vertigo, loss of balance, and
- 363 neuropsychiatric events such as headache, somnolence, and sleep disorders
- 364 (insomnia, abnormal dreams). These are usually mild and may decrease
- despite continued use.
- 366 Occasionally, more severe neuropsychiatric disorders have been reported such
- as: sensory and motor neuropathies (including paresthesia, tremor and ataxia),
- 368 convulsions, agitation or restlessness, anxiety, depression, mood changes,
- panic attacks, forgetfulness, confusion, hallucinations, aggression, psychotic
- or paranoid reactions and encephalopathy. Rare cases of suicidal ideation and
- 371 suicide have been reported though no relationship to drug administration has
- been confirmed.

Other infrequent adverse events include:

- 374 Cardiovascular Disorders: circulatory disturbances (hypotension,
- 375 hypertension, flushing, syncope), chest pain, tachycardia or palpitation,
- bradycardia, irregular pulse, extrasystoles, A-V block, and other transient
- 377 cardiac conduction alterations
- 378 Skin Disorders: rash, exanthema, erythema, urticaria, pruritus, edema, hair
- loss, erythema multiforme, and Stevens-Johnson syndrome
- 380 Musculoskeletal Disorders: muscle weakness, muscle cramps, myalgia, and
- 381 arthralgia

- 382 Respiratory Disorders: dyspnea, pneumonitis of possible allergic etiology 383 Other Symptoms: visual disturbances, vestibular disorders including tinnitus 384 and hearing impairment, asthenia, malaise, fatigue, fever, sweating, chills, 385 dyspepsia and loss of appetite 386 Laboratory 387 The most frequently observed laboratory alterations which could be possibly 388 attributable to drug administration were decreased hematocrit, transient 389 elevation of transaminases, leukopenia and thrombocytopenia. These 390 alterations were observed in patients with acute malaria who received 391 treatment doses of the drug and were attributed to the disease itself. 392 During prophylactic administration of mefloquine to indigenous populations 393 in malaria-endemic areas, the following occasional alterations in laboratory 394 values were observed: transient elevation of transaminases, leukocytosis or 395 thrombocytopenia. 396 Because of the long half-life of mefloquine, adverse reactions to Lariam may 397 occur or persist up to several weeks after the last dose. 398 **OVERDOSAGE** 399 **Symptoms and Signs** 400 In cases of overdosage with Lariam, the symptoms mentioned under 401 **ADVERSE REACTIONS** may be more pronounced. 402 **Treatment** 403 Patients should be managed by symptomatic and supportive care following 404 Lariam overdose. There are no specific antidotes. Monitor cardiac function (if 405 possible by ECG) and neuropsychiatric status for at least 24 hours. Provide 406 symptomatic and intensive supportive treatment as required, particularly for 407 cardiovascular disturbances. **DOSAGE AND ADMINISTRATION (see INDICATIONS AND USAGE)** 408 409 **Adult Patients** 410 Treatment of mild to moderate malaria in adults caused by P. vivax or
- 411 mefloquine-susceptible strains of *P. falciparum*
- 412 Five tablets (1250 mg) mefloquine hydrochloride to be given as a single oral
- dose. The drug should not be taken on an empty stomach and should be
- administered with at least 8 oz (240 mL) of water.
- 415 If a full-treatment course with Lariam does not lead to improvement within 48
- 416 to 72 hours, Lariam should not be used for retreatment. An alternative therapy
- should be used. Similarly, if previous prophylaxis with mefloquine has failed,
- 418 Lariam should not be used for curative treatment.

- Note: Patients with acute *P. vivax* malaria, treated with Lariam, are at high risk of relapse because Lariam does not eliminate exoerythrocytic (hepatic phase) parasites. To avoid relapse after initial treatment of the acute infection with Lariam, patients should subsequently be treated with an 8-aminoquinoline derivative (eg, primaquine).
- 424 Malaria Prophylaxis
- 425 One 250 mg Lariam tablet once weekly.
- 426 Prophylactic drug administration should begin 1 week before arrival in an
- 427 endemic area. Subsequent weekly doses should be taken regularly, always on
- 428 the same day of each week, preferably after the main meal. To reduce the risk
- of malaria after leaving an endemic area, prophylaxis must be continued for 4
- 430 additional weeks to ensure suppressive blood levels of the drug when
- 431 merozoites emerge from the liver. Tablets should not be taken on an empty
- stomach and should be administered with at least 8 oz (240 mL) of water.
- In certain cases, eg, when a traveler is taking other medication, it may be
- desirable to start prophylaxis 2 to 3 weeks prior to departure, in order to
- ensure that the combination of drugs is well tolerated (see **PRECAUTIONS**:
- 436 **Drug Interactions**).
- When prophylaxis with Lariam fails, physicians should carefully evaluate
- which antimalarial to use for therapy.

439 **Pediatric Patients**

- Treatment of mild to moderate malaria in pediatric patients caused by
- 441 mefloquine-susceptible strains of *P. falciparum*
- Twenty (20) to 25 mg/kg body weight. Splitting the total therapeutic dose into
- 2 doses taken 6 to 8 hours apart may reduce the occurrence or severity of
- adverse effects. Experience with Lariam in infants less than 3 months old or
- 445 weighing less than 5 kg is limited. The drug should not be taken on an empty
- stomach and should be administered with ample water. The tablets may be
- crushed and suspended in a small amount of water, milk or other beverage for
- administration to small children and other persons unable to swallow them
- 449 whole.
- 450 If a full-treatment course with Lariam does not lead to improvement within 48
- 451 to 72 hours, Lariam should not be used for retreatment. An alternative therapy
- should be used. Similarly, if previous prophylaxis with mefloquine has failed,
- Lariam should not be used for curative treatment.
- 454 In pediatric patients, the administration of Lariam for the treatment of malaria
- has been associated with early vomiting. In some cases, early vomiting has
- 456 been cited as a possible cause of treatment failure (see **PRECAUTIONS**). If a
- significant loss of drug product is observed or suspected because of vomiting,
- a second full dose of Lariam should be administered to patients who vomit

- less than 30 minutes after receiving the drug. If vomiting occurs 30 to 60
- 460 minutes after a dose, an additional half-dose should be given. If vomiting
- 461 recurs, the patient should be monitored closely and alternative malaria
- 462 treatment considered if improvement is not observed within a reasonable
- 463 period of time.
- The safety and effectiveness of Lariam to treat malaria in pediatric patients
- below the age of 6 months have not been established.
- 466 Malaria Prophylaxis
- 467 The following doses have been extrapolated from the recommended adult
- dose. Neither the pharmacokinetics, nor the clinical efficacy of these doses
- 469 have been determined in children owing to the difficulty of acquiring this
- 470 information in pediatric subjects. The recommended prophylactic dose of
- 471 Lariam is approximately 5 mg/kg body weight once weekly. One 250 mg
- Lariam tablet should be taken once weekly in pediatric patients weighing over
- 473 45 kg. In pediatric patients weighing less than 45 kg, the weekly dose
- decreases in proportion to body weight:
- 475 30 to 45 kg: 3/4 tablet
- 476 20 to 30 kg: 1/2 tablet
- 477 10 to 20 kg: 1/4 tablet
- 478 5 to 10 kg: 1/8 tablet*
- 479 *Approximate tablet fraction based on a dosage of 5 mg/kg body weight.
- Exact doses for children weighing less than 10 kg may best be prepared and
- 481 dispensed by pharmacists.
- 482 Experience with Lariam in infants less than 3 months old or weighing less
- than 5 kg is limited.

484 **HOW SUPPLIED**

- 485 Lariam is available as scored, white, round tablets, containing 250 mg of
- 486 mefloquine hydrochloride in unit-dose packages of 25 (NDC 0004-0172-02).
- 487 Imprint on tablets: LARIAM 250 ROCHE
- Tablets should be stored at 25°C (77°F); excursions permitted to 15° to 30°C
- 489 (59° to 86°F).

490 ANIMAL TOXICOLOGY

- 491 Ocular lesions were observed in rats fed mefloquine daily for 2 years. All
- 492 surviving rats given 30 mg/kg/day had ocular lesions in both eyes
- 493 characterized by retinal degeneration, opacity of the lens, and retinal edema.
- 494 Similar but less severe lesions were observed in 80% of female and 22% of
- male rats fed 12.5 mg/kg/day for 2 years. At doses of 5 mg/kg/day, only
- 496 corneal lesions were observed. They occurred in 9% of rats studied.

- 497 Revised: Month Year
- 498 **MEDICATION GUIDE**
- 499 This Medication Guide is intended only for travelers who are taking
- Lariam to prevent malaria. The information may not apply to patients who
- are sick with malaria and who are taking Lariam to treat malaria.
- An information wallet card is provided with this Medication Guide. Carry it
- with you when you are taking Lariam.
- This Medication Guide was revised in May 2004. Please read it before you
- start taking Lariam and each time you get a refill. There may be new
- 506 information. This Medication Guide does not take the place of talking with
- 507 your prescriber (doctor or other health care provider) about Lariam and
- malaria prevention. Only you and your prescriber can decide if Lariam is right
- 509 for you. If you cannot take Lariam, you may be able to take a different
- 510 medicine to prevent malaria.

What is the most important information I should know about Lariam?

- 512 1. Take Lariam exactly as prescribed to prevent malaria.
- Malaria is an infection that can cause death and is spread to humans
- through mosquito bites. If you travel to parts of the world where the
- mosquitoes carry the malaria parasite, you must take a malaria prevention
- medicine. Lariam is one of a small number of medications approved to
- prevent and to treat malaria. If taken correctly, Lariam is effective at
- preventing malaria but, like all medications, it may produce side effects in
- some patients.
- 520 2. Lariam can rarely cause serious mental problems in some patients.
- The most frequently reported side effects with Lariam, such as nausea,
- difficulty sleeping, and bad dreams are usually mild and do not cause
- people to stop taking the medicine. However, people taking Lariam
- occasionally experience severe anxiety, feelings that people are against
- 525 them, hallucinations (seeing or hearing things that are not there, for
- example), depression, unusual behavior, or feeling disoriented. There have
- been reports that in some patients these side effects continue after Lariam
- 528 is stopped. Some patients taking Lariam think about killing themselves,
- and there have been rare reports of suicides. It is not known whether
- Lariam was responsible for these suicides.
- 531 3. You need to take malaria prevention medicine before you travel to a
- malaria area, while you are in a malaria area, and after you return
- from a malaria area.
- Medicines approved in the United States for malaria prevention include
- Lariam, doxycycline, atovaquone/proguanil, hydroxychloroquine, and
- chloroquine. Not all of these drugs work equally as well in all areas of the

- 537 world where there is malaria. The chloroquines, for example, do not work 538 in areas where the malaria parasite has developed resistance to 539 chloroquine. Lariam may be effective against malaria that is resistant to 540 chloroquine or other drugs. All drugs to treat malaria have side effects that 541 are different for each one. For example, some may make your skin more 542 sensitive to sunlight (Lariam does not do this). However, if you use 543 Lariam to prevent malaria and you develop a sudden onset of anxiety, 544 depression, restlessness, confusion (possible signs of more serious mental 545 problems), or you develop other serious side effects, contact a doctor or 546 other health care provider. It may be necessary to stop taking Lariam and 547 use another malaria prevention medicine instead. If you can't get another 548 medicine, leave the malaria area. However, be aware that leaving the 549 malaria area may not protect you from getting malaria. You still need to 550 take a malaria prevention medicine.
- Who should not take Lariam?
- Do not take Lariam to **prevent** malaria if you
- have depression or had depression recently
- have had recent mental illness or problems, including anxiety disorder, schizophrenia (a severe type of mental illness), or psychosis (losing touch with reality)
- have or had seizures (epilepsy or convulsions)
- are allergic to quinine or quinidine (medicines related to Lariam)
- Tell your prescriber about all your medical conditions. Lariam may not be right for you if you have certain conditions, especially the ones listed below:
- **Heart disease.** Lariam may not be right for you.
- **Pregnancy.** Tell your prescriber if you are pregnant or plan to become pregnant. It is dangerous for the mother and for the unborn baby (fetus) to get malaria during pregnancy. Therefore, ask your prescriber if you should take Lariam or another medicine to prevent malaria while you are pregnant.
- **Breast-feeding.** Lariam can pass through your milk and may harm the baby. Therefore, ask your prescriber whether you will need to stop breast-feeding or use another medicine.
- Liver problems.
- 571 Tell your prescriber about all the medicines you take, including
- 572 prescription and non-prescription medicines, vitamins, and herbal
- 573 **supplements.** Some medicines may give you a higher chance of having
- serious side effects from Lariam.

- 575 How should I take Lariam?
- 576 Take Lariam exactly as prescribed. If you are an adult or pediatric
- patient weighing 45 kg (99 pounds) or less, your prescriber will tell you
- 578 the correct dose based on your weight.
- 579 To prevent malaria
- For adults and pediatric patients weighing over 45 kg, take 1 tablet of
 Lariam at least 1 week before you travel to a malaria area (or 2 to 3 weeks
 before you travel to a malaria area, if instructed by your prescriber). This
 starts the prevention and also helps you see how Lariam affects you and
 the other medicines you take. **Take 1 Lariam tablet once a week**, on the
 same day each week, while in a malaria area.
- Continue taking Lariam for 4 weeks after returning from a malaria area. If you cannot continue taking Lariam due to side effects or for other reasons, contact your prescriber.
- Take Lariam just after a meal and with at least 1 cup (8 ounces) of water.
- For children, Lariam can be given with water or crushed and mixed with
 water or sugar water. The prescriber will tell you the correct dose for
 children based on the child's weight.
- 593 If you are told by a doctor or other health care provider to stop taking 594 Lariam due to side effects or for other reasons, it will be necessary to take 595 another malaria medicine. You must take **malaria prevention medicine** 596 before you travel to a malaria area, while you are in a malaria area, and after you return from a malaria area. If you don't have access to 597 a doctor or other health care provider or to another medicine besides 598 599 Lariam and have to stop taking it, leave the malaria area. However, be aware that leaving the malaria area may not protect you from getting 600 malaria. You still need to take a malaria prevention medicine. 601
- 602 What should I avoid while taking Lariam?
- Halofantrine (marketed under various brand names), a medicine used to treat malaria. Taking both of these medicines together can cause serious heart problems that can cause death.
- **Do not become pregnant.** Women should use effective birth control while taking Lariam.
- Quinine, quinidine, or chloroquine (other medicines used to treat malaria). Taking these medicines with Lariam could cause changes in your heart rate or increase the risk of seizures.

611 **In addition:**

- **Be careful driving or in other activities** needing alertness and careful movements (fine motor coordination). Lariam can cause dizziness or loss of balance, even after you stop taking it.
- Be aware that certain vaccines may not work if given while you are
 taking Lariam. Your prescriber may want you to finish taking your
 vaccines at least 3 days before starting Lariam.

What are the possible side effects of Lariam?

- 619 Lariam, like all medicines, may cause side effects in some patients. The most
- 620 frequently reported side effects with Lariam when used for prevention of
- malaria include nausea, vomiting, diarrhea, dizziness, difficulty sleeping, and
- bad dreams. These are usually mild and do not cause people to stop taking the
- 623 medicine.
- Lariam may cause serious mental problems in some patients (see "What is the
- most important information I should know about Lariam?").
- 626 Lariam may affect your liver and your eyes if you take it for a long time. Your
- prescriber will tell you if you should have your eyes and liver checked while
- 628 taking Lariam.

What else should I know about preventing malaria?

- Find out whether you need malaria prevention. Before you travel, talk
- with your prescriber about your travel plans to determine whether you
- need to take medicine to prevent malaria. Even in those countries where
- malaria is present, there may be areas of the country that are free of
- malaria. In general, malaria is more common in rural (country) areas than
- in big cities, and it is more common during rainy seasons, when
- mosquitoes are most common. You can get information about the areas of
- the world where malaria occurs from the Centers for Disease Control and
- Prevention (CDC) and from local authorities in the countries you visit. If
- possible, plan your travel to reduce the risk of malaria.
- **Take medicine to prevent malaria infection**. Without malaria prevention
- medicine, you have a higher risk of getting malaria. Malaria starts with
- flu-like symptoms, such as chills, fever, muscle pains, and headaches.
- However, malaria can make you very sick or cause death if you don't seek
- medical help immediately. These symptoms may disappear for a while,
- and you may think you are well. But, the symptoms return later and then it
- may be too late for successful treatment.
- Malaria can cause confusion, coma, and seizures. It can cause kidney
- failure, breathing problems, and severe damage to red blood cells.
- However, malaria can be easily diagnosed with a blood test, and if
- caught in time, can be effectively treated.

651 652 653 654	If you get flu-like symptoms (chills, fever, muscle pains, or headaches) after you return from a malaria area, get medical help right away and tell your prescriber that you may have been exposed to malaria.
655 656 657 658	People who have lived for many years in areas with malaria may have some immunity to malaria (they do not get it as easily) and may not take malaria prevention medicine. This does not mean that you don't need to take malaria prevention medicine.
659 660 661 662 663 664 665	• Protect against mosquito bites . Medicines do not always completely prevent your catching malaria from mosquito bites. So protect yourself very well against mosquitoes. Cover your skin with long sleeves and long pants, and use mosquito repellent and bednets while in malaria areas. If you are out in the bush, you may want to pre-wash your clothes with permethrin. This is a mosquito repellent that may be effective for weeks after use. Ask your prescriber for other ways to protect yourself.
666	General information about the safe and effective use of Lariam.
667 668 669 670 671 672 673	Medicines are sometimes prescribed for conditions not listed in Medication Guides. If you have any concerns about Lariam, ask your prescriber. This Medication Guide contains certain important information for travelers visiting areas with malaria. Your prescriber or pharmacist can give you information about Lariam that was written for health care professionals. Do not use Lariam for a condition for which it was not prescribed. Do not share Lariam with other people.
674 675	This Medication Guide has been approved by the U.S. Food and Drug Administration.
676	Medication Guide Revised: May 2004
677	
678	Reprint of information wallet card:

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Lariam® (mefloquine hydrochloride) Tablets

Carry this information wallet card with you when you are taking Lariam.

You need to take malaria prevention medicine before you travel to a malaria area, while you are in a malaria area, and after you return from a malaria area.

If taken correctly, Lariam is effective at preventing malaria but, like all medications, it may produce side effects in some patients.

If you use Lariam to prevent malaria and you develop a sudden onset of anxiety, depression, restlessness, confusion (possible signs of more serious mental problems), or you develop other serious side effects, contact a doctor or other health care provider. It may be necessary to stop taking Lariam and use another malaria prevention medicine instead.

Other medicines approved in the United States for malaria prevention include: doxycycline, atovaquone/proguanil, hydroxychloroquine, and chloroquine. Not all malaria medicines work equally well in malaria areas. The chloroquines, for example, do not work in many parts of the world. If you can't get another medicine, leave the malaria area. However, be aware that leaving the malaria area may not protect you from getting malaria. You still need to take a malaria prevention medicine.

Please read the Medication Guide for additional information on Lariam.

Card Revised: May 2004

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682 Manufactured by: F. HOFFMANN-LA ROCHE LTD 683 684

Basel, Switzerland

Distributed by:

Roche

Pharmaceuticals

Roche Laboratories Inc. 340 Kingsland Street Nutley, New Jersey 07110-1199

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