AMERGE®
(naratriptan hydrochloride)
Tablets

DESCRIPTION
AMERGE Tablets contain naratriptan as the hydrochloride, which is a selective
5-hydroxytryptamine_1 receptor subtype agonist. Naratriptan hydrochloride is chemically
designated as N-methyl-3-(1-methyl-4-piperidinyl)-1H-indole-5-ethansulfonamide
monohydrochloride, and it has the following structure:

The empirical formula is C_{17}H_{25}N_{3}O_{2}S\cdot HCl, representing a molecular weight of 371.93.
Naratriptan hydrochloride is a white to pale yellow powder that is readily soluble in water. Each
AMERGE Tablet for oral administration contains 1.11 or 2.78 mg of naratriptan hydrochloride
equivalent to 1 or 2.5 mg of naratriptan, respectively. Each tablet also contains the inactive
ingredients croscarmellose sodium; hypromellose; lactose; magnesium stearate; microcrystalline
cellulose; triacetin; and titanium dioxide, iron oxide yellow (2.5-mg tablet only), and indigo
carmine aluminum lake (FD&C Blue No. 2) (2.5-mg tablet only) for coloring.

CLINICAL PHARMACOLOGY
Mechanism of Action: Naratriptan binds with high affinity to 5-HT_{1D} and 5-HT_{1B} receptors
and has no significant affinity or pharmacological activity at 5-HT_{2-4} receptor subtypes or at
adrenergic \(\alpha_1, \alpha_2,\) or \(\beta;\) dopaminergic \(D_1\) or \(D_2;\) muscarinic; or benzodiazepine receptors.
The therapeutic activity of naratriptan in migraine is generally attributed to its agonist activity
at 5-HT_{1D/1B} receptors. Two current theories have been proposed to explain the efficacy of
5-HT_{1D/1B} receptor agonists in migraine. One theory suggests that activation of 5-HT_{1D/1B}
receptors located on intracranial blood vessels, including those on the arteriovenous
anastomoses, leads to vasoconstriction, which is correlated with the relief of migraine headache.
The other hypothesis suggests that activation of 5-HT_{1D/1B} receptors on sensory nerve endings in
the trigeminal system results in the inhibition of pro-inflammatory neuropeptide release.
In the anesthetized dog, naratriptan has been shown to reduce the carotid arterial blood flow
with little or no effect on arterial blood pressure or total peripheral resistance. While the effect on
blood flow was selective for the carotid arterial bed, increases in vascular resistance of up to
30% were seen in the coronary arterial bed. Naratriptan has also been shown to inhibit trigeminal
nerve activity in rat and cat. In 10 human subjects with suspected coronary artery disease (CAD) undergoing coronary artery catheterization, there was a 1% to 10% reduction in coronary artery diameter following subcutaneous injection of 1.5 mg of naratriptan.

**Pharmacokinetics:** Naratriptan tablets are well absorbed, with about 70% oral bioavailability. Following administration of a 2.5-mg tablet orally, the peak concentrations are obtained in 2 to 3 hours. After administration of 1- or 2.5-mg tablets, the $C_{\text{max}}$ is somewhat (about 50%) higher in women (not corrected for milligram-per-kilogram dose) than in men. During a migraine attack, absorption was slower, with a $T_{\text{max}}$ of 3 to 4 hours. Food does not affect the pharmacokinetics of naratriptan. Naratriptan displays linear kinetics over the therapeutic dose range.

The steady-state volume of distribution of naratriptan is 170 L. Plasma protein binding is 28% to 31% over the concentration range of 50 to 1,000 ng/mL.

Naratriptan is predominantly eliminated in urine, with 50% of the dose recovered unchanged and 30% as metabolites in urine. In vitro, naratriptan is metabolized by a wide range of cytochrome P450 isoenzymes into a number of inactive metabolites.

The mean elimination half-life of naratriptan is 6 hours. The systemic clearance of naratriptan is 6.6 mL/min/kg. The renal clearance (220 mL/min) exceeds glomerular filtration rate, indicating active tubular secretion. Repeat administration of naratriptan tablets does not result in drug accumulation.

**Special Populations:**

**Age:** A small decrease in clearance (approximately 26%) was observed in healthy elderly subjects (65 to 77 years) compared to younger patients, resulting in slightly higher exposure (see PRECAUTIONS).

**Race:** The effect of race on the pharmacokinetics of naratriptan has not been examined.

**Renal Impairment:** Clearance of naratriptan was reduced by 50% in patients with moderate renal impairment (creatinine clearance, 18 to 39 mL/min) compared to the normal group. Decrease in clearances resulted in an increase of mean half-life from 6 hours (healthy) to 11 hours (range, 7 to 20 hours). The mean $C_{\text{max}}$ increased by approximately 40%. The effects of severe renal impairment (creatinine clearance, $\leq 15$ mL/min) on the pharmacokinetics of naratriptan has not been assessed (see CONTRAINDICATIONS and DOSAGE AND ADMINISTRATION).

**Hepatic Impairment:** Clearance of naratriptan was decreased by 30% in patients with moderate hepatic impairment (Child-Pugh grade A or B). This resulted in an approximately 40% increase in the half-life (range, 8 to 16 hours). The effects of severe hepatic impairment (Child-Pugh grade C) on the pharmacokinetics of naratriptan have not been assessed (see CONTRAINDICATIONS and DOSAGE AND ADMINISTRATION).

**Drug Interactions:** In normal volunteers, coadministration of single doses of naratriptan tablets and alcohol did not result in substantial modification of naratriptan pharmacokinetic parameters.

From population pharmacokinetic analyses, coadministration of naratriptan and fluoxetine, beta-blockers, or tricyclic antidepressants did not affect the clearance of naratriptan.
Naratriptan does not inhibit monoamine oxidase (MAO) enzymes and is a poor inhibitor of P450; metabolic interactions between naratriptan and drugs metabolized by P450 or MAO are therefore unlikely.

**Oral Contraceptives:** Oral contraceptives reduced clearance by 32% and volume of distribution by 22%, resulting in slightly higher concentrations of naratriptan. Hormone replacement therapy had no effect on pharmacokinetics in older female patients.

Smoking increased the clearance of naratriptan by 30%.

**CLINICAL TRIALS**

The efficacy of AMERGE Tablets in the acute treatment of migraine headaches was evaluated in 6 randomized, double-blind, placebo-controlled studies of which 4 used the recommended dosing regimen and were conducted as outpatient trials. Three of these studies enrolled adult patients who were predominantly female (86%) and Caucasian (96%) with a mean age of 41 (range, 18 to 65). One study enrolled adolescents with a mean age of 14 (range, 12 to 17). In the adolescent study, 54% of the patients were female and 89% were Caucasian. In all studies, patients were instructed to treat at least 1 moderate to severe headache. Headache response, defined as a reduction in headache severity from moderate or severe pain to mild or no pain, was assessed up to 4 hours after dosing. Associated symptoms such as nausea, vomiting, photophobia, and phonophobia were also assessed. Maintenance of response was assessed for up to 24 hours postdose. A second dose of AMERGE Tablets or other medication was allowed 4 to 24 hours after the initial treatment for recurrent headache. The frequency and time to use of these additional treatments were also determined.

In all 3 trials in adults utilizing the recommended dosage regimen and outpatient use, the percentage of patients achieving headache response 4 hours after treatment, the primary outcome measure, was significantly greater among patients receiving AMERGE compared to those who received placebo. In all studies, response to 2.5 mg was numerically greater than response to 1 mg and in the largest of the 3 studies, there was a statistically significant greater percentage of patients with headache response at 4 hours in the 2.5-mg group compared to the 1-mg group. The results are summarized in Table 1.

<table>
<thead>
<tr>
<th>Study</th>
<th>Placebo</th>
<th>AMERGE 1.0 mg</th>
<th>AMERGE 2.5 mg</th>
</tr>
</thead>
<tbody>
<tr>
<td>Study 1</td>
<td>34% (n = 122)</td>
<td>50%* (n = 117)</td>
<td>60%* (n = 127)</td>
</tr>
<tr>
<td>Study 2</td>
<td>27% (n = 104)</td>
<td>52%* (n = 208)</td>
<td>66%*† (n = 199)</td>
</tr>
<tr>
<td>Study 3</td>
<td>32% (n = 169)</td>
<td>54%* (n = 166)</td>
<td>65%* (n = 167)</td>
</tr>
</tbody>
</table>

*p<0.05 in comparison with placebo.

†p<0.05 in comparison with 1 mg.
In the single study in adolescents, there were no statistically significant differences between any of the treatment groups. The headache response rates at 4 hours (n) were 65% (n = 74), 67% (n = 78), and 64% (n = 70) for placebo, 1-mg, and 2.5-mg groups, respectively.

Comparisons of drug performance based upon results obtained in different clinical trials are never reliable. Because studies are conducted at different times, with different samples of patients, by different investigators, employing different criteria and/or different interpretations of the same criteria, under different conditions (dose, dosing regimen, etc.), quantitative estimates of treatment response and the timing of response may be expected to vary considerably from study to study.

The estimated probability of achieving an initial headache response in adults over the 4 hours following treatment is depicted in Figure 1.

Figure 1. Estimated Probability of Achieving Initial Headache Response Within 4 Hours*

* The figure shows the probability over time of obtaining headache response (no or mild pain) following treatment with AMERGE Tablets. The averages displayed are based on pooled data from the 3 controlled clinical trials providing evidence of efficacy (Studies 1, 2, and 3). In this Kaplan-Meier plot, patients not achieving response within 240 minutes were censored at 240 minutes.
For patients with migraine-associated nausea, photophobia, and phonophobia at baseline, there was a lower incidence of these symptoms 4 hours following administration of 1- and 2.5-mg AMERGE Tablets compared to placebo.

Four to 24 hours following the initial dose of study treatment, patients were allowed to use additional treatment for pain relief in the form of a second dose of study treatment or other medication. The estimated probability of patients taking a second dose or other medication for migraine over the 24 hours following the initial dose of study treatment is summarized in Figure 2.

**Figure 2. Estimated Probability of Patients Taking a Second Dose of AMERGE Tablets or Other Medication for Migraine Over the 24 Hours Following the Initial Dose of Study Treatment**

* Kaplan-Meier plot based on data obtained in the 3 controlled clinical trials (Studies 1, 2, and 3) providing evidence of efficacy with patients not using additional treatments censored at 24 hours. The plot also includes patients who had no response to the initial dose.

Remedication was discouraged prior to 4 hours postdose.

There is no evidence that doses of 5 mg provide a greater effect than 2.5 mg. There was no evidence to suggest that treatment with AMERGE was associated with an increase in the severity or frequency of migraine attacks. The efficacy of AMERGE Tablets was unaffected by presence
of aura; gender, age, or weight of the patient; oral contraceptive use; or concomitant use of common migraine prophylactic drugs (e.g., beta-blockers, calcium channel blockers, tricyclic antidepressants). There was insufficient data to assess the impact of race on efficacy.

INDICATIONS AND USAGE

AMERGE Tablets are indicated for the acute treatment of migraine attacks with or without aura in adults. AMERGE Tablets are not intended for the prophylactic therapy of migraine or for use in the management of hemiplegic or basilar migraine (see CONTRAINDICATIONS). Safety and effectiveness of AMERGE Tablets have not been established for cluster headache, which is present in an older, predominantly male population.

CONTRAINDICATIONS

AMERGE Tablets should not be given to patients with history, symptoms, or signs of ischemic cardiac, cerebrovascular, or peripheral vascular syndromes. In addition, patients with other significant underlying cardiovascular diseases should not receive AMERGE Tablets. Ischemic cardiac syndromes include, but are not limited to, angina pectoris of any type (e.g., stable angina of effort and vasospastic forms of angina such as the Prinzmetal variant), all forms of myocardial infarction, and silent myocardial ischemia. Cerebrovascular syndromes include, but are not limited to, strokes of any type as well as transient ischemic attacks. Peripheral vascular disease includes, but is not limited to, ischemic bowel disease (see WARNINGS).

Because AMERGE Tablets may increase blood pressure, they should not be given to patients with uncontrolled hypertension (see WARNINGS).

AMERGE Tablets are contraindicated in patients with severe renal impairment (creatinine clearance, <15 mL/min) (see CLINICAL PHARMACOLOGY and DOSAGE AND ADMINISTRATION).

AMERGE Tablets are contraindicated in patients with severe hepatic impairment (Child-Pugh grade C) (see CLINICAL PHARMACOLOGY and DOSAGE AND ADMINISTRATION).

AMERGE Tablets should not be administered to patients with hemiplegic or basilar migraine.

AMERGE Tablets should not be used within 24 hours of treatment with another 5-HT1 agonist, an ergotamine-containing or ergot-type medication like dihydroergotamine or methysergide.

AMERGE Tablets are contraindicated in patients with hypersensitivity to naratriptan or any of the components.

WARNINGS

AMERGE Tablets should only be used where a clear diagnosis of migraine has been established.
Risk of Myocardial Ischemia and/or Infarction and Other Adverse Cardiac Events: Because of the potential of this class of compounds (5-HT\textsubscript{1B/1D} agonists) to cause coronary vasospasm, naratriptan should not be given to patients with documented ischemic or vasospastic coronary artery disease (CAD) (see CONTRAINDICATIONS). It is strongly recommended that 5-HT\textsubscript{1} agonists (including naratriptan) not be given to patients in whom unrecognized CAD is predicted by the presence of risk factors (e.g., hypertension, hypercholesterolemia, smoker, obesity, diabetes, strong family history of CAD, female with surgical or physiological menopause, or male over 40 years of age) unless a cardiovascular evaluation provides satisfactory clinical evidence that the patient is reasonably free of coronary artery and ischemic myocardial disease or other significant underlying cardiovascular disease. The sensitivity of cardiac diagnostic procedures to detect cardiovascular disease or predisposition to coronary artery vasospasm is modest, at best. If, during the cardiovascular evaluation, the patient’s medical history, electrocardiographic, or other investigations reveal findings indicative of, or consistent with, coronary artery vasospasm or myocardial ischemia, naratriptan should not be administered (see CONTRAINDICATIONS).

For patients with risk factors predictive of CAD, who are determined to have a satisfactory cardiovascular evaluation, it is strongly recommended that administration of the first dose of naratriptan take place in the setting of a physician’s office or similar medically staffed and equipped facility. Because cardiac ischemia can occur in the absence of clinical symptoms, consideration should be given to obtaining on the first occasion of use an electrocardiogram (ECG) during the interval immediately following administration of AMERGE Tablets, in these patients with risk factors.

It is recommended that patients who are intermittent long-term users of 5-HT\textsubscript{1} agonists, including AMERGE Tablets, and who have or acquire risk factors predictive of CAD, as described above, undergo periodic cardiovascular evaluation as they continue to use AMERGE Tablets.

The systematic approach described above is intended to reduce the likelihood that patients with unrecognized cardiovascular disease will be inadvertently exposed to naratriptan.

Cardiac Events and Fatalities Associated With 5-HT\textsubscript{1} Agonists: Naratriptan can cause coronary artery vasospasm (see CLINICAL PHARMACOLOGY). Serious adverse cardiac events, including acute myocardial infarction, life-threatening disturbances of cardiac rhythm, and death have been reported within a few hours following the administration of 5-HT\textsubscript{1} agonists. Considering the extent of use of 5-HT\textsubscript{1} agonists in patients with migraine, the incidence of these events is extremely low.

Premarking Experience With AMERGE Tablets: Among approximately 3,500 patients with migraine who participated in premarketing clinical trials of naratriptan tablets, 4 patients treated with single oral doses of naratriptan ranging from 1 to 10 mg experienced
asymptomatic ischemic ECG changes with at least 1, who took 7.5 mg, likely due to coronary
vasospasm.

Cerebrovascular Events and Fatalities With 5-HT\textsubscript{1} Agonists: Cerebral hemorrhage,
subarachnoid hemorrhage, stroke, and other cerebrovascular events have been reported in
patients treated with 5-HT\textsubscript{1} agonists, and some have resulted in fatalities. In a number of cases, it
appears possible that the cerebrovascular events were primary, the agonist having been
administered in the incorrect belief that the symptoms experienced were a consequence of
migraine, when they were not. It should be noted that patients with migraine may be at increased
risk of certain cerebrovascular events (e.g., stroke, hemorrhage, transient ischemic attack).

Other Vasospasm-Related Events: 5-HT\textsubscript{1} agonists may cause vasospastic reactions other
than coronary artery spasm. Both peripheral vascular ischemia and colonic ischemia with
abdominal pain and bloody diarrhea have been reported with naratriptan.

Serotonin Syndrome: The development of a potentially life-threatening serotonin syndrome
may occur with triptans, including treatment with AMERGE, particularly during combined use
with selective serotonin reuptake inhibitors (SSRIs) or serotonin norepinephrine reuptake
inhibitors (SNRIs). If concomitant treatment with naratriptan and an SSRI (e.g., fluoxetine,
paroxetine, sertraline, fluvoxamine, citalopram, escitalopram) or SNRI (e.g., venlafaxine,
duloxetine) is clinically warranted, careful observation of the patient is advised, particularly
during treatment initiation and dose increases. Serotonin syndrome symptoms may include
mental status changes (e.g., agitation, hallucinations, coma), autonomic instability (e.g.,
thachycardia, labile blood pressure, hyperthermia), neuromuscular aberrations (e.g., hyperreflexia,
incoordination), and/or gastrointestinal symptoms (e.g., nausea, vomiting, diarrhea).

Increase in Blood Pressure: In healthy volunteers, dose-related increases in systemic blood
pressure have been observed after administration of up to 20 mg of oral naratriptan. At the
recommended doses, the elevations are generally small, although an increase of systolic pressure
of 32 mmHg was seen in 1 patient following a single 2.5-mg dose. The effect may be more
pronounced in the elderly and hypertensive patients. A patient who was mildly hypertensive (the
baseline blood pressure was 150/98) experienced a significant increase in blood pressure to
204/144 mmHg 225 minutes after administration of a 10-mg oral dose. Significant elevation in
blood pressure, including hypertensive crisis, has been reported on rare occasions in patients
receiving 5-HT\textsubscript{1} agonists with and without a history of hypertension. Naratriptan is
contraindicated in patients with uncontrolled hypertension (see CONTRAINDICATIONS).

An 18% increase in mean pulmonary artery pressure and an 8% increase in mean aortic
pressure was seen following dosing with 1.5 mg of subcutaneous naratriptan in a study
evaluating 10 subjects with suspected CAD undergoing cardiac catheterization.

Hypersensitivity: Hypersensitivity (anaphylaxis/anaphylactoid) reactions may occur in
patients receiving naratriptan. Such reactions can be life threatening or fatal. In general,
hypersensitivity reactions to drugs are more likely to occur in individuals with a history of
sensitivity to multiple allergens (see CONTRAINDICATIONS).
PRECAUTIONS

General: Chest discomfort (including pain, pressure, heaviness, tightness) has been reported after administration of 5-HT\textsubscript{1} agonists, including AMERGE Tablets. These events have not been associated with arrhythmias or ischemic ECG changes in clinical trials with AMERGE Tablets. Because naratriptan may cause coronary artery vasospasm, patients who experience signs or symptoms suggestive of angina following naratriptan should be evaluated for the presence of CAD or a predisposition to Prinzmetal variant angina before receiving additional doses of naratriptan, and should be monitored electrocardiographically if dosing is resumed and similar symptoms recur. Similarly, patients who experience other symptoms or signs suggestive of decreased arterial flow, such as ischemic bowel syndrome or Raynaud syndrome following naratriptan administration should be evaluated for atherosclerosis or predisposition to vasospasm (see CONTRAINDICATIONS and WARNINGS).

AMERGE Tablets should also be administered with caution to patients with diseases that may alter the absorption, metabolism, or excretion of drugs, such as impaired renal or hepatic function (see CLINICAL PHARMACOLOGY, CONTRAINDICATIONS, and DOSAGE AND ADMINISTRATION).

Care should be taken to exclude other potentially serious neurological conditions before treating headache in patients not previously diagnosed with migraine or who experience a headache that is atypical for them. There have been rare reports where patients received 5-HT\textsubscript{1} agonists for severe headaches that were subsequently shown to have been secondary to an evolving neurologic lesion (see WARNINGS).

For a given attack, if a patient has no response to the first dose of AMERGE, the diagnosis of migraine should be reconsidered before administration of a second dose.

Binding to Melanin-Containing Tissues: In rats treated with a single oral dose (10 mg/kg) of radiolabeled naratriptan, the elimination half-life of radioactivity from the eye was 90 days, suggesting that naratriptan and/or its metabolites may bind to the melanin of the eye. Because there could be accumulation in melanin-rich tissues over time, this raises the possibility that naratriptan could cause toxicity in these tissues after extended use. Although no systematic monitoring of ophthalmologic function was undertaken in clinical trials, and no specific recommendations for ophthalmologic monitoring are offered, prescribers should be aware of the possibility of long-term ophthalmologic effects.

Changes in the Precorneal Tear Film: Dogs receiving oral naratriptan showed transient changes in the precorneal tear film. Corneal stippling was seen at the lowest dose tested, 1 mg/kg/day, and occurred intermittently from day 1 throughout the first 2 to 3 weeks of treatment. Although a no-effect dose was not established, the exposure at the lowest dose tested was approximately 5 times the human exposure after a 5-mg oral dose.

Information for Patients: See PATIENT INFORMATION at the end of this labeling for the text of the separate leaflet provided for patients.

Patients should be cautioned about the risk of serotonin syndrome with the use of naratriptan or other triptans, especially during combined use with SSRIs or SNRIs.
**Laboratory Tests:** No specific laboratory tests are recommended for monitoring patients prior to and/or after treatment with AMERGE Tablets.

**Drug Interactions:**

**Selective Serotonin Reuptake Inhibitors/Serotonin Norepinephrine Reuptake Inhibitors and Serotonin Syndrome:** Cases of life-threatening serotonin syndrome have been reported during combined use of SSRIs or SNRIs and triptans (see WARNINGS).

**Ergot-Containing Drugs:** Ergot-containing drugs have been reported to cause prolonged vasospastic reactions. Because there is a theoretical basis that these effects may be additive, use of ergotamine-containing or ergot-type medications (like dihydroergotamine or methysergide) and naratriptan within 24 hours is contraindicated (see CONTRAINDICATIONS).

**Other 5-HT1 Agonists:** The administration of naratriptan with other 5-HT1 agonists has not been evaluated in migraine patients. Because their vasospastic effects may be additive, coadministration of naratriptan and other 5-HT1 agonists within 24 hours of each other is not recommended (see CONTRAINDICATIONS).

**Drug/Laboratory Test Interactions:** AMERGE Tablets are not known to interfere with commonly employed clinical laboratory tests.

**Carcinogenesis, Mutagenesis, Impairment of Fertility:**

**Carcinogenesis:** Lifetime carcinogenicity studies, 104 weeks in duration, were carried out in mice and rats by oral gavage. There was no evidence of an increase in tumors related to naratriptan administration in mice receiving up to 200 mg/kg/day. That dose was associated with a plasma area-under-the-curve (AUC) exposure that was 110 times the exposure in humans receiving the maximum recommended daily dose of 5 mg. Two rat studies were conducted, 1 using a standard diet and the other a nitrite-supplemented diet (naratriptan can be nitrosated in vitro to form a mutagenic product that has been detected in the stomachs of rats fed a high nitrite diet). Doses of 5, 20, and 90 mg/kg were associated with week 13 AUC exposures that in the standard diet study were 7, 40, and 236 times, respectively, and in the nitrite-supplemented diet study were 7, 29, and 180 times, respectively, the exposure attained in humans given the maximum recommended daily dose of 5 mg. In both studies, there was an increase in the incidence of thyroid follicular hyperplasia in high-dose males and females and in thyroid follicular adenomas in high-dose males. In the standard diet study only, there was also an increase in the incidence of benign c-cell adenomas in the thyroid of high-dose males and females. The exposures achieved at the no-effect dose for thyroid tumors were 40 (standard diet) and 29 (nitrite-supplemented diet) times the exposure achieved in humans receiving the maximum recommended daily dose of 5 mg. In the nitrite-supplemented diet study only, the incidence of benign lymphocytic thymoma was increased in all treated groups of females. It was not determined if the nitrosated product is systemically absorbed. However, no changes were seen in the stomachs of rats in that study.

**Mutagenesis:** Naratriptan was not mutagenic when tested in 2 gene mutation assays, the Ames test and the in vitro thymidine locus mouse lymphoma assay. It was not clastogenic in 2 cytogenetics assays, the in vitro human lymphocyte assay and the in vivo mouse micronucleus assay.
assay. Naratriptan can be nitrosated in vitro to form a mutagenic product (WHO nitrosation assay) that has been detected in the stomachs of rats fed a nitrite-supplemented diet.

**Impairment of Fertility:** In a reproductive toxicity study in which male and female rats were dosed prior to and throughout the mating period with 10, 60, 170, or 340 mg/kg/day (plasma exposures [AUC] approximately 11, 70, 230, and 470 times, respectively, the human exposure at the maximum recommended daily dose [MRDD] of 5 mg), there was a treatment-related decrease in the number of females exhibiting normal estrous cycles at doses of 170 mg/kg/day or greater and an increase in preimplantation loss at 60 mg/kg/day or greater. In high-dose group males, testicular/epididymal atrophy accompanied by spermatozoa depletion reduced mating success and may have contributed to the observed preimplantation loss. The exposures achieved at the no-effect doses for preimplantation loss, anestrus, and testicular effects were approximately 11, 70, and 230 times, respectively, the exposures in humans receiving the MRDD.

In a study in which rats were dosed orally with 10, 60, or 340 mg/kg/day for 6 months, changes in the female reproductive tract including atrophic or cystic ovaries and anestrus were seen at the high dose. The exposure at the no-effect dose of 60 mg/kg was approximately 85 times the exposure in humans receiving the MRDD.

**Pregnancy:** Pregnancy Category C. There are no adequate and well-controlled studies in pregnant women; therefore, naratriptan should be used during pregnancy only if the potential benefit justifies the potential risk to the fetus.

To monitor fetal outcomes of pregnant women exposed to AMERGE, GlaxoSmithKline maintains a Naratriptan Pregnancy Registry. Healthcare providers are encouraged to register patients by calling (800) 336-2176.

In reproductive toxicity studies in rats and rabbits, oral administration of naratriptan was associated with developmental toxicity (embryolethality, fetal abnormalities, pup mortality, offspring growth retardation) at doses producing maternal plasma drug exposures as low as 11 and 2.5 times, respectively, the exposure in humans receiving the MRDD of 5 mg.

When pregnant rats were administered naratriptan during the period of organogenesis at doses of 10, 60, or 340 mg/kg/day, there was a dose-related increase in embryonic death, with a statistically significant difference at the highest dose, and incidences of fetal structural variations (incomplete/irregular ossification of skull bones, sternebrae, ribs) were increased at all doses. The maternal plasma exposures (AUC) at these doses were approximately 11, 70, and 470 times the exposure in humans at the MRDD. The high dose was maternally toxic, as evidenced by decreased maternal body weight gain during gestation. A no-effect dose for developmental toxicity in rats exposed during organogenesis was not established.

When doses of 1, 5, or 30 mg/kg/day were given to pregnant Dutch rabbits throughout organogenesis, the incidence of a specific fetal skeletal malformation (fused sternebrae) was increased at the high dose, and increased incidences of embryonic death and fetal variations (major blood vessel variations, supernumerary ribs, incomplete skeletal ossification) were observed at all doses (4, 20, and 120 times, respectively, the MRDD on a body surface area
Maternal toxicity (decreased body weight gain) was evident at the high dose in this study. In a similar study in New Zealand White rabbits (1, 5, or 30 mg/kg/day throughout organogenesis), decreased fetal weights and increased incidences of fetal skeletal variations were observed at all doses (maternal exposures equivalent to 2.5, 19, and 140 times exposure in humans receiving the MRDD), while maternal body weight gain was reduced at 5 mg/kg or greater. A no-effect dose for developmental toxicity in rabbits exposed during organogenesis was not established.

When female rats were treated with 10, 60, or 340 mg/kg/day during late gestation and lactation, offspring behavioral impairment (tremors) and decreased offspring viability and growth were observed at doses of 60 mg/kg or greater, while maternal toxicity occurred only at the highest dose. Maternal exposures at the no-effect dose for developmental effects in this study were approximately 11 times the exposure in humans receiving the MRDD.

**Nursing Mothers:** Naratriptan-related material is excreted in the milk of rats. Therefore, caution should be exercised when considering the administration of AMERGE Tablets to a nursing woman.

**Pediatric Use:** Safety and effectiveness of AMERGE Tablets in pediatric patients (younger than 18 years) have not been established.

One randomized, placebo-controlled clinical trial evaluating oral naratriptan (0.25 to 2.5 mg) in pediatric patients aged 12 to 17 years evaluated a total of 300 adolescent migraineurs. This study did not establish the efficacy of oral naratriptan compared to placebo in the treatment of migraine in adolescents (see CLINICAL TRIALS). Adverse events observed in this clinical trial were similar in nature to those reported in clinical trials in adults.

**Geriatric Use:** The use of AMERGE Tablets in elderly patients is not recommended.

Naratriptan is known to be substantially excreted by the kidney, and the risk of adverse reactions to this drug may be greater in elderly patients who have reduced renal function. In addition, elderly patients are more likely to have decreased hepatic function; they are at higher risk for CAD; and blood pressure increases may be more pronounced in the elderly. Clinical studies of AMERGE Tablets did not include patients over 65 years of age.

**ADVERSE REACTIONS**

Serious cardiac events, including some that have been fatal, have occurred following the use of 5-HT1 agonists. These events are extremely rare and most have been reported in patients with risk factors predictive of CAD. Events reported have included coronary artery vasospasm, transient myocardial ischemia, myocardial infarction, ventricular tachycardia, and ventricular fibrillation (see CONTRAINDICATIONS, WARNINGS, and PRECAUTIONS).

**Incidence in Controlled Clinical Trials:** The most common adverse events were paresthesias, dizziness, drowsiness, malaise/fatigue, and throat/neck symptoms, which occurred at a rate of 2% and at least 2 times placebo rate. Since patients treated only 1 to 3 headaches in the controlled clinical trials, the opportunity for discontinuation of therapy in response to an
adverse event was limited. In a long-term, open-label study where patients were allowed to treat multiple migraine attacks for up to 1 year, 15 patients (3.6%) discontinued treatment due to adverse events.

Table 2 lists adverse events that occurred in 5 placebo-controlled clinical trials of approximately 1,752 exposures to placebo and AMERGE Tablets in adult migraine patients. The events cited reflect experience gained under closely monitored conditions of clinical trials in a highly selected patient population. In actual clinical practice or in other clinical trials, these frequency estimates may not apply, as the conditions of use, reporting behavior, and the kinds of patients treated may differ. Only events that occurred at a frequency of 2% or more in the group treated with AMERGE Tablets 2.5 mg and were more frequent in that group than in the placebo group are included in Table 2. From this table, it appears that many of these adverse events are dose related.

Table 2. Treatment-Emergent Adverse Events Reported by at Least 2% of Patients in Placebo-Controlled Migraine Trials

<table>
<thead>
<tr>
<th>Adverse Event Type</th>
<th>Placebo (n = 498)</th>
<th>AMERGE 1 mg (n = 627)</th>
<th>AMERGE 2.5 mg (n = 627)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Atypical sensation</td>
<td>1%</td>
<td>2%</td>
<td>4%</td>
</tr>
<tr>
<td>Paresthesias (all types)</td>
<td>&lt;1%</td>
<td>1%</td>
<td>2%</td>
</tr>
<tr>
<td>Gastrointestinal</td>
<td>5%</td>
<td>6%</td>
<td>7%</td>
</tr>
<tr>
<td>Nausea</td>
<td>4%</td>
<td>4%</td>
<td>5%</td>
</tr>
<tr>
<td>Neurological</td>
<td>3%</td>
<td>4%</td>
<td>7%</td>
</tr>
<tr>
<td>Dizziness</td>
<td>1%</td>
<td>1%</td>
<td>2%</td>
</tr>
<tr>
<td>Drowsiness</td>
<td>&lt;1%</td>
<td>1%</td>
<td>2%</td>
</tr>
<tr>
<td>Malaise/fatigue</td>
<td>1%</td>
<td>2%</td>
<td>2%</td>
</tr>
<tr>
<td>Pain and pressure sensation</td>
<td>2%</td>
<td>2%</td>
<td>4%</td>
</tr>
<tr>
<td>Throat/neck symptoms</td>
<td>1%</td>
<td>1%</td>
<td>2%</td>
</tr>
</tbody>
</table>

One event (vomiting) present in more than 1% of patients receiving AMERGE Tablets occurred more frequently on placebo than on naratriptan 2.5 mg.

AMERGE Tablets are generally well tolerated. Most adverse reactions were mild and transient.

The incidence of adverse events in placebo-controlled clinical trials was not affected by age or weight of the patients, duration of headache prior to treatment, presence of aura, use of prophylactic medications, or tobacco use. There was insufficient data to assess the impact of race on the incidence of adverse events.

Other Events Observed in Association With the Administration of AMERGE Tablets: In the paragraphs that follow, the frequencies of less commonly reported adverse clinical events are presented. Because the reports include events observed in open and uncontrolled studies, the role of AMERGE Tablets in their causation cannot be reliably...
determined. Furthermore, variability associated with adverse event reporting, the terminology
used to describe adverse events, etc., limit the value of the quantitative frequency estimates
provided. Event frequencies are calculated as the number of patients reporting an event divided
by the total number of patients (n = 3,557) exposed to oral naratriptan doses up to 10 mg. All
reported events are included except those already listed in the previous table, those too general to
be informative, and those not reasonably associated with the use of the drug. Events are further
classified within body system categories and enumerated in order of decreasing frequency using
the following definitions: frequent adverse events are those occurring in at least 1/100 patients,
infrequent adverse events are those occurring in 1/100 to 1/1,000 patients, and rare adverse
events are those occurring in fewer than 1/1,000 patients.

**Atypical Sensations:** Frequent were warm/cold temperature sensations. Infrequent were
feeling strange and burning/stinging sensation.

**Cardiovascular:** Infrequent were palpitations, increased blood pressure, tachyarrhythmias,
and abnormal ECG (PR prolongation, QTc prolongation, ST/T wave abnormalities, premature
ventricular contractions, atrial flutter, or atrial fibrillation), and syncope. Rare were bradycardia,
varicosities, hypotension, and heart murmurs.

**Ear, Nose, and Throat:** Frequent were ear, nose, and throat infections. Infrequent were
phonophobia, sinusitis, upper respiratory inflammation, and tinnitus. Rare were allergic rhinitis;
labyrinthitis; ear, nose, and throat hemorrhage; and hearing difficulty.

**Endocrine and Metabolic:** Infrequent were thirst and polydipsia, dehydration, and fluid
retention. Rare were hyperlipidemia, hypercholesterolemia, hypothyroidism, hyperglycemia,
glycosuria and ketonuria, and parathyroid neoplasm.

**Eye:** Frequent was photophobia. Infrequent was blurred vision. Rare were eye pain and
discomfort, sensation of eye pressure, eye hemorrhage, dry eyes, difficulty focusing, and
scotoma.

**Gastrointestinal:** Frequent were hyposalivation and vomiting. Infrequent were dyspeptic
symptoms, diarrhea, gastrointestinal discomfort and pain, gastroenteritis, and constipation. Rare
were abnormal liver function tests, abnormal bilirubin levels, hemorrhoids, gastritis, esophagitis,
salivary gland inflammation, oral itching and irritation, regurgitation and reflux, and gastric
ulcers.

**Hematological Disorders:** Infrequent was increased white cells. Rare were
thrombocytopenia, quantitative red cell or hemoglobin defects, anemia, and purpura.

**Lower Respiratory Tract:** Infrequent were bronchitis, cough, and pneumonia. Rare were
tracheitis, asthma, pleuritis, and airway constriction and obstruction.

**Musculoskeletal:** Infrequent were muscle pain, arthralgia and articular rheumatism, muscle
cramps and spasms, joint and muscle stiffness, tightness, and rigidity. Rare were bone and
skeletal pain.

**Neurological:** Frequent was vertigo. Infrequent were tremors, cognitive function disorders,
sleep disorders, and disorders of equilibrium. Rare were compressed nerve syndromes,
confusion, sedation, hyperesthesia, coordination disorders, paralysis of cranial nerves, decreased
consciousness, dreams, altered sense of taste, neuralgia, neuritis, aphasia, hypoesthesia, motor
retardation, muscle twitching and fasciculation, psychomotor restlessness, and convulsions.

**Non-Site Specific:** Infrequent were chills and/or fever, descriptions of odor or taste, edema
and swelling, allergies, and allergic reactions. Rare were spasms and mobility disorders.

**Pain and Pressure Sensations:** Frequent were pressure/tightness/heaviness sensations.

**Psychiatry:** Infrequent were anxiety, depressive disorders, and detachment. Rare were
agression and hostility, agitation, hallucinations, panic, and hyperactivity.

**Reproduction:** Rare were lumps of female reproductive tract, breast inflammation,
inflammation of vagina, inflammation of fallopian tube, breast discharge, endometrium
disorders, decreased libido, and lumps of breast.

**Skin:** Infrequent were sweating, skin rashes, pruritus, and urticaria. Rare were skin erythema,
dermatitis and dermatosis, hair loss and alopecia, pruritic skin rashes, acne and folliculitis,
allergic skin reactions, macular skin/rashes, skin photosensitivity, photodermatitis, skin flakiness,
and dry skin.

**Urology:** Infrequent were bladder inflammation and polyuria and diuresis. Rare were urinary
tract hemorrhage, urinary urgency, pyelitis, and urinary incontinence.

**Observed During Clinical Practice:** The following section enumerates potentially important
adverse events that have occurred in clinical practice and that have been reported spontaneously
to various surveillance systems. The events enumerated represent reports arising from both
domestic and nondomestic use of naratriptan. These events do not include those already listed in
the ADVERSE REACTIONS section above. Because the reports cite events reported
spontaneously from worldwide postmarketing experience, frequency of events and the role of
naratriptan in their causation cannot be reliably determined.

**Cardiovascular:** Angina, myocardial infarction (see WARNINGS).

**Gastrointestinal:** Colonic ischemia (see WARNINGS).

**Lower Respiratory:** Dyspnea.

**Miscellaneous:** Hypersensitivity, including anaphylaxis/anaphylactoid reactions, in some
cases severe (e.g., circulatory collapse) (see WARNINGS).

**Neurologic:** Cerebral vascular accident, including transient ischemic attack, subarachnoid
hemorrhage, and cerebral infarction (see WARNINGS); serotonin syndrome.

**DRUG ABUSE AND DEPENDENCE**

In one clinical study enrolling 12 subjects, all of whom had experience using oral opiates and
other psychoactive drugs, AMERGE Tablets produced less intense subjective responses
ordinarily associated with many drugs of abuse than did codeine (30 to 90 mg).

**OVERDOSAGE**

A patient who was mildly hypertensive experienced a significant increase in blood pressure
after administration of a 10-mg dose starting at 30 minutes (baseline value of 150/98 to
204/144 mmHg 225 minutes). This event resolved after treatment with antihypertensive therapy.
Oral administration of 25 mg of naratriptan in 1 healthy young male subject increased blood
pressure from 120/67 mmHg pretreatment up to 191/113 mmHg at approximately 6 hours postdose and resulted in adverse events including lightheadedness, tension in the neck, tiredness, and loss of coordination. Blood pressure returned to near baseline by 8 hours after dosing without any pharmacological intervention.

Another subject experienced asymptomatic ischemic ECG changes likely due to coronary artery vasospasm approximately 2 hours following a 7.5-mg oral dose.

The elimination half-life of naratriptan is about 6 hours (see CLINICAL PHARMACOLOGY), and therefore monitoring of patients after overdose with AMERGE Tablets should continue for at least 24 hours or while symptoms or signs persist. There is no specific antidote to naratriptan. Standard supportive treatment should be applied as required. If the patient presents with chest pain or other symptoms consistent with angina pectoris, ECG monitoring should be performed for evidence of ischemia. It is unknown what effect hemodialysis or peritoneal dialysis has on the serum concentrations of naratriptan.

**DOSAGE AND ADMINISTRATION**

In controlled clinical trials, single doses of 1 and 2.5 mg of AMERGE Tablets taken with fluid were effective for the acute treatment of migraines in adults. A greater proportion of patients had headache response following a 2.5-mg dose than following a 1-mg dose (see CLINICAL TRIALS). Individuals may vary in response to doses of AMERGE Tablets. The choice of dose should therefore be made on an individual basis, weighing the possible benefit of the 2.5-mg dose with the potential for a greater risk of adverse events. If the headache returns or if the patient has only partial response, the dose may be repeated once after 4 hours, for a maximum dose of 5 mg in a 24-hour period. There is evidence that doses of 5 mg do not provide a greater effect than 2.5 mg.

The safety of treating, on average, more than 4 headaches in a 30-day period has not been established.

**Renal Impairment:** The use of AMERGE is contraindicated in patients with severe renal impairment (creatinine clearance, <15 mL/min) because of decreased clearance of the drug (see CONTRAINDICATIONS and CLINICAL PHARMACOLOGY). In patients with mild to moderate renal impairment, the maximum daily dose should not exceed 2.5 mg over a 24-hour period and a lower starting dose should be considered.

**Hepatic Impairment:** The use of AMERGE is contraindicated in patients with severe hepatic impairment (Child-Pugh grade C) because of decreased clearance (see CONTRAINDICATIONS and CLINICAL PHARMACOLOGY). In patients with mild or moderate hepatic impairment, the maximum daily dose should not exceed 2.5 mg over a 24-hour period and a lower starting dose should be considered (see CLINICAL PHARMACOLOGY).

**HOW SUPPLIED**

AMERGE Tablets 1 and 2.5 mg of naratriptan (base) as the hydrochloride. AMERGE Tablets, 1 mg, are white, D-shaped, film-coated tablets debossed with “GX CE3” on one side in blister packs of 9 tablets (NDC 0173-0561-00). AMERGE Tablets, 2.5 mg, are green, D-shaped,
film-coated tablets debossed with “GX CE5” on one side in blister packs of 9 tablets (NDC 0173-0562-00).

Store at controlled room temperature, 20° to 25°C (68° to 77°F) (see USP).

PATIENT INFORMATION

The following wording is contained in a separate leaflet provided for patients.

Information for the Patient
AMERGE® (naratriptan hydrochloride) Tablets

Please read this leaflet carefully before you take AMERGE Tablets. This leaflet provides a summary of the information available about your medicine. Please do not throw away this leaflet until you have finished your medicine. You may need to read this leaflet again. This leaflet does not contain all the information on AMERGE Tablets. For further information or advice, ask your doctor or pharmacist.

Information About Your Medicine:

The name of your medicine is AMERGE (naratriptan hydrochloride) Tablets. It can be obtained only by prescription from your doctor. The decision to use AMERGE Tablets is one that you and your doctor should make jointly, taking into account your individual preferences and medical circumstances. If you have risk factors for heart disease (such as high blood pressure, high cholesterol, obesity, diabetes, smoking, strong family history of heart disease, or you are postmenopausal or a male over 40), you should tell your doctor, who should evaluate you for heart disease in order to determine if AMERGE is appropriate for you. The majority of those who have taken AMERGE Tablets have not experienced any significant side effects. Rarely, deaths and/or serious heart problems have been reported with this class of medicines; in all but a few instances, however, these deaths and/or serious heart problems occurred in people with heart disease and it was not clear whether these medicines were a contributing factor.

1. The Purpose of Your Medicine:

AMERGE Tablets are intended to relieve your migraine, but not to prevent or reduce the number of attacks you experience. Use AMERGE Tablets only to treat an actual migraine attack.

2. Important Questions to Consider Before Taking AMERGE Tablets:

If the answer to any of the following questions is YES or if you do not know the answer, then please discuss it with your doctor before you use AMERGE Tablets.

• Are you pregnant? Do you think you might be pregnant? Are you trying to become pregnant? Are you not using adequate contraception? Are you breastfeeding?

• Do you have any chest pain, heart disease, shortness of breath, or irregular heartbeats? Have you had a heart attack?

• Do you have risk factors for heart disease (such as high blood pressure, high cholesterol, obesity, diabetes, smoking, strong family history of heart disease, or you are postmenopausal or a male over 40)?
• Have you had a stroke, transient ischemic attacks (TIAs), or Raynaud syndrome?
• Do you have high blood pressure?
• Have you ever had to stop taking this or any other medicine because of an allergy or other problems?
• Are you taking any other migraine medicines, including other 5-HT₁ agonists such as IMITREX® (sumatriptan/sumatriptan succinate), or medicines containing ergotamine, dihydroergotamine, or methysergide?
• Are you taking any medicine for depression or other disorders such as selective serotonin reuptake inhibitors (SSRIs) or serotonin norepinephrine reuptake inhibitors (SNRIs)?
  Common SSRIs are citalopram HBr (CELEXA®), escitalopram oxalate (LEXAPRO®), paroxetine (PAXIL®), fluoxetine (PROZAC®/SARAFEM®), olanzapine/fluoxetine (SYMBYAX®), sertraline (ZOLOFT®), and fluvoxamine. Common SNRIs are duloxetine (CYMBALTA®) and venlafaxine (EFFEXOR®).*
• Have you had, or do you have, any disease of the kidney or liver?
• Is this headache different from your usual migraine attacks?

Remember, if you answered YES to any of the above questions, then discuss it with your doctor.

3. The Use of AMERGE Tablets During Pregnancy:
Do not use AMERGE Tablets if you are pregnant, think you might be pregnant, are trying to become pregnant, or are not using adequate contraception, unless you have discussed this with your doctor.

4. How to Use AMERGE Tablets:
For adults, the usual dose is a single tablet taken whole with fluids. It may be given at any time after the headache starts. For an individual attack, if you have no response to the first tablet, do not take a second tablet without first talking to your doctor. If you need more relief due to a partial response or return of your headache after the first tablet, a second tablet may be taken, but not sooner than 4 hours following the first tablet. Do not take more than a total of 2 AMERGE Tablets in any 24-hour period. If you have kidney or liver disease, take as directed by your doctor.

5. Side Effects to Watch for:
• Some patients experience pain or tightness in the chest or throat when using AMERGE Tablets. If this happens to you, then discuss it with your doctor before using any more AMERGE Tablets. If the chest pain, tightness, or pressure is severe or does not go away, call your doctor immediately.
• If you have sudden and/or severe abdominal pain following AMERGE Tablets, call your doctor immediately.
• Some people may have a reaction called serotonin syndrome when they use certain types of antidepressants, SSRIs or SNRIs, while taking AMERGE Tablets. Symptoms may include confusion, hallucinations, fast heart beat, feeling faint, fever, sweating, muscle spasm,
difficulty walking, and/or diarrhea. Call your doctor immediately if you have any of these symptoms after taking AMERGE Tablets.

- Shortness of breath; wheeziness; heart throbbing, swelling of eyelids, face, or lips; or a skin rash, skin lumps, or hives happens rarely. If it happens to you, then tell your doctor immediately. Do not take any more AMERGE Tablets unless your doctor tells you to do so.

- Some people may have feelings of tingling, heat, flushing (redness of face lasting a short time), heaviness or pressure after treatment with AMERGE Tablets. A few people may feel drowsy, dizzy, tired, or sick. Tell your doctor of these symptoms at your next visit.

- If you feel unwell in any other way or have any symptoms that you do not understand, you should contact your doctor immediately.

6. What to Do if an Overdose Is Taken:

If you have taken more medicine than you have been told, contact either your doctor, hospital emergency department, or nearest poison control center immediately.

7. Storing Your Medicine:

Keep your medicine in a safe place where children cannot reach it. It may be harmful to children. Store your medicine away from heat and light. Do not store at temperatures above 77°F (25°C). If your medicine has expired (the expiration date is printed on the treatment pack), throw it away as instructed. If your doctor decides to stop your treatment, do not keep any leftover medicine unless your doctor tells you to. Throw away your medicine as instructed.

*AMERGE, IMITREX, and PAXIL are registered trademarks of GlaxoSmithKline. The other brands listed are trademarks of their respective owners and are not trademarks of GlaxoSmithKline. The makers of these brands are not affiliated with and do not endorse GlaxoSmithKline or its products.