PRODUCT INFORMATION

Significant hypertensive episodes, including hypertensive crises, have been reported on rare occasions in patients with ar without a history of hypertension (see WARNINGS), incidence in Controlled Clinical Triats: Among 3,653 pa-tients treated with IMITREX Nasal Spray in active- and on the section of the section of the section of the section.

placebo-controlled clinical trials, less than 0.4% of natients pinebo-controlled clinical trials, less than 0.4% of patients withdrew for reasons related to adverse events. Table 2 lists adverse ovents that occurred in worldwide placebo-con-trolled clinical trials in 3,419 migraineurs. 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 froncy estimates may not apply, as the conditions of use, orting behavior, and the kinds of patients treated may

Only events that occurred at a frequency of 1% or more in the IMITREX Nasal Spray 20-mg treatment group and were more frequent in that group than in the placebo group are included in Table 2.

nophobia also occurred in more than 1% of patients but

Phonophobia also occurres in most state of the second state of the events in controlled clinical trials was not affected by gen-der, weight, or age of the patients; use of prophysicatic medi-cations; or presence of aurs. There were insufficient data to assess the impact of race on the incidence of adverse events. Other Events Observed in Association With the Adminis-tration of IMITREX Nasal Spray: In the paragraphs that follow, the frequencies of less commonly reported adverse clinical events are presented. Because the reports include events abserved in one and uncontrolled studies. events observed in open and uncontrolled studies, the role of IMITREX Nasal Spray in their causation cannot be reliably determined. Furthermore, variability associated with addetermined. Furthermore, variability associated with ad-verse event reporting, the terminology used to describe ad-verse events, etc., limit the value of the quantitative frequency estimates provided. Event frequencies are calcu-lated as the number of patients who used IMITREX Nasal lated as the number of patients who used IMITREX Nasai Spray (5, 10, or 20 mg in controlled and uncontrolled trials) and reported an event divided by the total number of pa-tients (n = 3,711) exposed to IMITREX Nasai Spray. All re-ported 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. previous table, those too general to be use of the drug. those not reasonably associated within body system categories and enumerated in order of decreasing frequency using the following definitions: 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. occurring in 1/100 to 1/1,000 patients and rare adverse svents are those occurring in fewer than 1/1,000 patients. Atypical Sensations: Infrequent were tingling, warm/hot sensation, numbness, pressure sensation, feeling strange, feeling of heaviness, feeling of tightness, paresthesia, cold sensation, and tight feeling in head. Rare were dysesthesia and prickling sensation.

vascular: Infrequent were flushing and hyperten-ee WARNINGS), palpitations, tachycardis, changes and used what unot, parpitations, cachycardin, changes in ECG, and arrhythmia (see WARNINGS and PRECAU-TIONS). Rare were abdominal aortic aneurysm, hypoten-sion, bradycardin, pallor, and phlebitis. Chest Symptoms: Infrequent were chest tightness, chest disconfort, and chest pressure/heaviness (see PRECAU-TIONS). Comparison

TIONS: General

Ear, Nose, and Throat: Infrequent were disturbance of hearing and ear infection. Rare were otalgia and Meniere dise Endoceia

e and Metabolic: Infrequent was thirst. Rare galactorrhea, hypothyroidism, and weight loss. Infrequent were irritation of eyes and visual disturwere gr Eve:

ntestinal: Infrequent were abdominal discomfort, diarrhea, dysphagia, and gastroesophageal reflux. Rare were constipation, flatulence/eructation, hematemesis, in-testinal obstruction, melena, gastroenteritis, colitis, hemor-

rhage of gastrointestinal tract, and pancreatitis. Mouth and Teeth: Infrequent was disorder of mouth and tongue (e.g., burning of tongue, numbress of tongue, dry month)

mouth). Musculoskoletal: Infrequent were neck pain/stiffness, backache, weakness, joint symptoms, arthritis, and myal-gia. Rare were muscle cramps, tetany, intervertebral disc disorder, and muscle stiffness.

al: Infrequent were drowsiness/sedation, anxi-Neurological: infrequent were drowsness/sedation, ann-ety, aleep disturbances, tremors, syncope, ahivera, chills, de-pression, agitation, sensation of lightness, and mental con-fusion. Rare were difficulty concentrating, hunger, lacrima-tion, memory disturbances, monoplegia/diplegia, apathy, disturbance of smell, disturbance of emotions, dysarthria, facial pain, intoxication, stress, decreased appetite, diffi-able conditions and mediane of children of children of the stress and the conditions and mediane of children of children of the stress of the conditions and mediane of children of the stress of the stress of the conditions and mediane of children of the stress of the st culty coordinating, euphoria, and neoplasm of pituitary. Respiratory: Infrequent were dyspnea and lower respira-tory tract infection. Rare was asthma.

Skin: Infrequent were rash/skin eruption, pruritus, and erythema. Rare were herpes, swelling of face, sweating, and ling of skin tal: Infrequent were dysuria, disorder of breasts

and dysmenorrhea. Rare were endometriosis and increas

Miscellaneous: Infrequent were cough, edema, and fever. Rare were hypersensitivity, swelling of extremities, voice disturbances, difficulty in walking, and lymphadenopathy. Other Events Observed in the Clinical Development of IMITREX: The following adverse events occurred in clinical

Table 2. Treatment-Emergent Adverse Events Reported by at Least 1% of Patients in Controlled Migraine Trials

	the second se	Percent of Pat	ients Reporting	
Adverse Event Type	Placebo (n = 704)	IMITREX 5 mg (n = 496)	IMITREX 10 mg (n = 1007)	IMITREX 20 mg (n = 1212)
Atypical sensations Burning sensation	0.1%	0.4%	0.6%	1.4%
Ear, nose, and throat Disorder/discomfort of nasal cavity/sinuses Throat discomfort	2.4% 0.9%	2.8% 0.8%	2.5% 1.8%	3.8% 2.4%
Gastrointestinal Nausea and/or vomiting	11.3%	12.2%	11.0%	13.5%
Neurological Bad/unusual taste Dizziness/vertigo	1.7% 0.9%	13.5% 1.0%	19.3% 1.7%	24.5% 1.4%

trials with IMITREX Injection and IMITREX Tablets. Because the reports include ovents observed in open and un-controlled studies, the role of IMITREX in their causation cannot be reliably determined. All reported events are in-cluded except those already listed, those too general to be informative, and those not reasonably associated with the

informative, and those not reasonably associated with the use of the drug. Breasts: Breast swelling; cysts, lumps, and masses of breasts; nipple discharge; primary malignant breast neo-plasm; and tenderness. Cardiovascular: Abnormal pulse, angins, atherosclerosis, cerebral ischemia, corebrovascular lesion, heart block, pe-ripherai cyandrome, thromosis, pulsating sensations. Raynaud syndrome, thrombosis, transient myocardial ischemia, various tran-inst ECC hearter (corrections). issue ECG changes (nonspecific ST or T wave changes, pro-longation of PR or QT intervals, sinus arrhythmis, nonsu-tained ventricular premature beats, isolated junctional e-topic beats, atrial ectopic beats, delayed activation of the right ventricle) and w odilatio

Ear, Nose, and Throat: Allergic rhinitis; ear, nose, and throat hemorrhage; external otitis; feeling of fullness in the ear(s); hearing disturbances; hearing loss; nasal inflamma-tion; sensitivity to noise; sinusitis; tinnitus; and upper restion; se piratory inflam ation.

Endocri and Metabolic: Dehydration; endocrine cysts lumps, and masses; elevated thyrotropin stimulating hor-mone (TSH) levels; fluid disturbances; hyperglycemia; hypoglycemia; polydipsia; and weight gain. Eye: Accommodation disorders, blindness and low vision,

conjunctivitis, disorders of sclera, external ocular muscle disorders, eye edema and swelling, eye itching, eye hemor-rhage, eye pain, keratitis, mydriasis, and vision alterations. Gastrointestinal: Abdominal distention, dental pain, dis turbances of liver function tests, dyspetite symptoms, feel-ings of gastrointestinal pressure, gallstones; gastric symp-toms, gastriits, gastrointestinal pain, hypersalivation, hypo-salivation, oral itching and irritation, peptie ulcer, retching, salivary gland swelling, and swallowing disorders Hematological Disorders: Anemia. rs: Ane

Injection Site Reactio

Miscollaneous: Contusions, fluid retention, hematoma, hypersensitivity to various agents, jaw discomfort, miscella-neous laboratory abnormalities, overdose, "serotonin agoneous laboratory abnormalities, ov nist effect", and speech disturbance.

mast entert, and speech disturbance. Musculoskeletal: Acquired musculoskeletal deformity, ar-thraigia and articular rheumatitis, muscle ntrophy, muscle tiredness, musculoskeletal inflammation, need to flex calf muscles, rigidity, tightness, and various joint disturbances (pain, stiffness, swelling, ache). Neurological: Aggressiveness, bradylogia, cluster head-ache convulsione databaneat disturbances of used.

ache, convulsions, detachment, disturbances of taste, drug ahuse, dystonia, facial paralysis, globus hystericus, halluci-nations, headache, heat sensitivity, hyperesthesia, hysteria, ache, convulsio increased alertheses, mean sensitivity, hyperestinesia, hysteria, increased alertheses, malaise/fatigue, migraine, motor dys-function, mycelonia, neuralgia, neurotic disorders, paraly-sis, personality change, phobia, photophobia, psychomotor disorders, radiculopathy, raised intracranial pressure, re-laxation, stinging sensations, transient hemiplegia, simul-taneous hot and cold sensations, suicide, tickling sensa-tions, twitching and varning.

taneous not and cold sensations, suicide, tickling sensa-tions, twitching, and yawning. Pain and Other Pressure Sensations: Chest pain, neck tightness/pressure, throat/jaw pain/tightness/pressure, and pain (location specified). Respiratory: Breathing disorders, bronchitis, diseases of

Skin: Dry/scaly skin, eczema, seborrheic dermatitis, skin nodules, skin tenderness, tightness of skin, and wrinkling of

skin.

Urogenital: Abortion, abnormal menstrual cycle, bladder inflammation, hematuria, inflammation of fallopian tubes, intermentation, nematuria, innammation of failopian tubes, intermentstrual bleeding, menstruation symptoms, micturi-tion disorders, renal calculus, urethritis, urinary frequency, and urinary infections.

Postmarketing Experience (Reports for Subcutaneous or Oral Sumatriptan): The following section enumerates potentially important adverse events that have occurred clinical practice and that have been reported spontaneou to various surveillance systems. The events enumerated represent reports arising from both domestic and nonde mestic use of oral or subcutaneous dosage forms of sumatriptan. The events enumerated include all except those already listed in the ADVERSE REACTIONS secti

above or those too general to be informative. Because the reports cite events reported spontaneously from worldwide postmarketing experience, frequency of events and the role of sumatriptan in their causation cannot be reliably deterin mined, it is assumed, however, that systemic reactions fol-lowing sumatriptan use are likely to be similar regardless of route of administration.

Blood: Hemolytic anemia, pancytopenia, thrombocytopenin

Atrial fibrillation, cardiomyopathy, colo ischemia (see WARNINGS), Prinzmetal variant angina, pul-monary embolism, shock, thrombophlebitis. *Ear, Nose, and Throat*: Deafness.

Eye: Ischemic optic neuropathy, retinal artery occlusion, retinal vein thrombosis, loss of vision.

Gastrointestinal: Ischemic colitis with rectal bleeding (see WARNINGS), xerr Hepatic: Elevated liver function tests.

Neurological: Central nervous system vasculitis, cerebro-vascular accident, dysphäsia, subarachnoid hemorrhage. Non-Site Specific: Angioneurotic edema, cyanosis, death (see WARNINGS), temporal arteritis.

Psychiatry: Panic disorder. spiratory: Bronchospasm in patients with and without a history of asthma.

Skin: Exacehation of sunburn, hypersensitivity reactions (allergic vasculitis, erythema, pruritus, rash, shortness of breath, urticaria; in addition, severe anaphylaxis/anaphy-lactoid reactions have been reported [see WARNINGS]), photosensitivity.

Urogenital: Acute renal failure. DRUG ABUSE AND DEPENDENCE

One clinical study with IMITREX (sumatriptan succinate) Injection enrolling 12 patients with a history of substance abuse failed to induce subjective behavior and/or physio-logic response ordinarily associated with drugs that have an established potential for abuse.

OVERDOSAGE

In clinical trials, the highest single doses of IMITREX Nasal Spray administered without significant adverse effects were 40 mg to 12 volunteers and 40 mg to 85 migraine patients, which is twice the highest single recommended dose. In ad-dition, 12 volunteers were administered a total daily dose of 60 mg (20 mg 3 times daily) for 3.5 days without signific se eve

Overdose in animals has been fatal and has been heralded Overdose in animais has over least incitivity, ptosis, ery-by convulsions, tremor, paralysis, inactivity, ptosis, ery-thema of the extremities, abnormal respiration, cyanosis, thema of the extremities, and larrimation. The eliminathema of the extremities, abnormal respiration, cyanosis, ataxia, mydriasis, salivation, and lacrimation. The elimina-tion hall-life of sumatriptan is about 2 hours (see CLINI-CAL PHARMACOLOGY), and therefore monitoring of pa-tients after overdose with IMITREX Nasal Spray should them after overdose with IMITREX Nasal Spray should continue for at least 10 hours or while symptoms or signs persist. It is unknown what effect hemodialysis or perio-neal dialysis has on the serum concentrations of sumatrip-

DOSAGE AND ADMINISTRATION

DOSAGE AND ADMINISTRATION In controlled clinical trials, single doses of 5, 10, or 20 mg of IMITREX Nasal Spray administered into 1 nostril were ef-fective for the acute treatment of migraine in adults. A greater proportion of patients had headache response fol-lowing a 20-mg dose than following a 5- or 10-mg dose (see CLINICAL TRIALS). Individuals may vary in response to doses of IMITREX Nasal Spray. The choice of dose should therefore be made on an individual basis, weighing the pos-sible benefit of the 20-mg dose with the potential for a greater risk of adverse events. A 10-mg dose may be greater risk of adverse events. A 10-mg dose may be achieved by the administration of a single 5-mg dose in each nostril. There is evidence that doses above 20 mg do not pro-vide a greater effect than 20 mg.

Continued on next page

This product information is based on lebeling in effect on August 1, 2002. Further information is available at: GlaxoSmithKline, PO Box 13398, Research Triangle Park, NC 27709. 1-888-825-5249. Corporate Web Site: www.gsk.co

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Imitrex Nasal Spray-Cont.

If the headache returns, the dose may be repeated once af-ter 2 hours, not to exceed a total daily dose of 40 mg. The safety of treating an average of more than 4 headaches in a 30-day period has not been established. HOW SUPPLIED

IMITREX Nasal Spray 5 mg (NDC 0173-0524-00) and 20 mg (NDC 0173-0523-00) are each supplied in boxes of 6 ansal spray devices. Each unit does spray supplies 5 and 20 mg, respectively, of sumatriptan. Store between 36° and 86°F (2° and 30°C). Protect from light

ANIMAL TOXICOLOGY

Corneal Opacities: Dogs receiving oral sumatriptan devel-oped corneal opacities and defects in the corneal epithelium. Corneal opacities were seen at the lowest dosage tested. al opac 2 mg/kg per day, and were present after 1 month of treat-ment. Defects in the corneal epithelium were noted in a 60-week study. Earlier examinations for these toxicities were not conducted and no-effect doses were not established; however, the relative exposure at the lowest does tested was approximately 5 times the human exposure after a 100-mg oral dose or 3 times the human exposure after a 6-mg suboral cose of 3 times the human exposure after a being sub-cutaneous dose or 22 times the human exposure after a sin-gle 20-mg intranasal dose. There is evidence of alterations in corneal appearance on the first day of intranasal dosing to dogs. Changes were noted at the lowest dose tested, which was approximately 2 times the maximum single hu-man intranasal dose of 20 mg on a mg/m² basis.

PATIENT INFORMATION

The following wording is contained in a separate leaflet pro-vided for patients.

Information for the Patient IMITREX® (sumatriptan) Nasal Spray. Please read this leaflet carefully before you administer IMITREX Nasal Spray. This provides a summary of the in-formation available on 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 IMITREX Nasal Spray. For further information on IMITREX Nasal Spray. For further information or advice, ask your doctor or pharma

formation About Your Media

Information About Your Medicine: The name of your medicine is IMITREX (sumatriptan) Na-sal Spray. It can be obtained only by prescription from your doctor. The decision to use IMITREX Nasal Spray is one that you and your doctor should make jointly taking into account your individual preferences and medical circum-stances. If you have risk factors for heart disease (such as which bland account your individual preferences and medical circum-stances. If you have risk factors for heart disease (such as Account your individual preservences and mencal circum-stances. 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 doc-tor, who should evaluate you for heart disease in order to determine if IMITREX is appropriate for you. Although the vast majority of those who have taken IMITREX have not experienced any significant side effects, some individuals have been reported. In all but a few instances, however, se-rious problema occurred in people with known heart disease and it was not clear whether IMITREX was a contributory factor in these deaths. **1. The Purpose of Your Medicine:** IMITREX Nasal Spray is intended to relieve your migraine, but not to prevent or reduce the number of attacks you ex-perience. Use IMITREX Nasal Spray only to treat an actual migraine attack.

igraine atta

- migraine attack.
 Important Questions to Consider Before Using IMITREX Nasal Spray:
 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 IMITREX Nasal Spray.
 Are you pregnant? Do you think you might be pregnant? Are you pregnant? Are you using inadequate contraception? Are you breastfielding?
 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 disea blood pressure, high cholesterol, obesity, diabetes, smok-ing, strong family history of heart disease, or you are post-menopausal or a maje 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 medi-

DOCKET

- Tave you taking any other migraine medicines, including other 5-HT, agonists or any other medicines, including ergotamine, dihydroergotamine, or methysergide?
 Are you taking any medicine for depression (monoanine oxidase inhibitors or selective serotonin reuptake inhibit-tion 1958-10
- tors [SSRIs])?
- · Have you had, or do you have, any disease of the liver or kidney?
- Have, you had, or do you have, epilepsy or seizures? Is this headache different from your usual migraine at-tacks?
- Remember, if you answered YES to any of the above ques-tions, then discuss it with your doctor.

rmation will be superseded by supplements and subsequent edit

3. The Use of IMITREX Nasal Spray During Prognancy: Do not use IMITREX Nasal Spray if you are pregnant, think you might be pregnant, are trying to become pregnant, or are not using adequate contraception, unless you have dis-cussed this with your doctor.

4. How to Use IMITREX Nasal Spray:

Before using IMITREX Nasal Spray, see the instruction pamphlet accompanying the product. For adults, the usual dose is a single nasal spray administered into 1 nostril. If your headache comes back, a second nasal spray may be ad-ministered anytime after 2 hours of administering the first ministered anytime after 2 hours of administering the first spray. For any attack where you have no response to the first nasal spray, do not take a second nasal spray without first consulting with your doctor. Do not administer more than a total of 40 mg of IMITREX Nasal Spray in any 24-hour period. The effects of long-term repeated use of IMI-TREX Nasal Spray on the surfaces of the nose and throat hour north comparison studied. The offset of the statement have not been specifically studied. The safety of treating an average of more than 4 headaches in a 30-day period has not been established.

5. Side Effects to Watch for:

Side theory to watch nor: Some patients experience pain or tightness in the chest or throat when using IMITREX Nasal Spray. If this happens to you, then discuss it with your doctor before using any more IMITREX Nasal Spray. If the chest pain is severe or

does not go away, call your doctor immediately. • If you have sudden and/or severe abdominal pain follow-ing IMITREX Nasal Spray, call your doctor immediately. Shortness of breath; wheeziness; heart throbbing; swell-ing 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 IMITREX Na-

sal Spray 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 pres-sure after treatment with IMITREX Nasal Spray. A few

(redne people may feel drowsy, dizzy, tired, sick, or may experi-ence nasal irritation. 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 docdiately. tor in

tor immediately. 6. What to Do if an Overdose is Taken: If you have taken more medicines than you have been told, contact either your doctor, hospital emergency department, or nearest poison control center immediately. 7. Storing Your Medicine:

7. Storing Your Medicine: Keep your medicine in a safe place where children cannot reach it. It may be harmful to children. Store your medi-cines away from heat and light. Do not store at temp-eratures above 86°F (30°C), or below 36°F (2°C). If your medicine has expired (the expiration date is printed on the treatment pack), throw it away as instructed. If your doctor heiden the term of the same store of the same stor decides to stop your treatment, do not keep any leftover medicine unless your doctor tells you to. Throw away your medicine as instructed.

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1 2002/RL-1005 Shown in Product Identification Guide, page 316 April 2

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IMITREX® Im 'I-trex

umatriptan succinate)

DESCRIPTION

IMITREX Tablets contain sumstriptan (as the succinate), a selective 5-hydroxytryptamine, receptor subtype agonist. Sumatriptan succinate is chemically designated as 3-12-(dimethylamino)ethyll-N-methyl-indole-5-methanesulfonamide succinate (1:1).

mide succinate (1:1). The empirical formula is $C_{14}H_{21}N_3O_2S^+C_4H_8O_{4*}$ represent-ing a molecular weight of 413.5. Sumatriptan succinate is a white to off-white powder that is readily soluble in water and in saline. Each IMITREX Tables for oral administration contains 35, 70, or 140 mg of sumatriptan succinate equiv-alent to 25, 50, or 100 mg of sumatriptan, respectively. Each tablet also contains the inactive ingredients crossermellose sodium, iron oxide (100-mg tablet only), lactose, magnesium stearste microcrastiling cellulace and titanium disside stearate, microcrystalline cellulose, and titanium dioxide. CLINICAL PHARMACOLOGY

CLINICAL PHARMACOLOGY Mechanism of Action: Sumatriptan is an agonist for a vascular 5-hydroxytryptamine, receptor subtype (probably a member of the 5-HT₁₀ family) having only a weak affinity for 5-HT₁₄, b-HT₂₆, and 6-HT₂ receptors and no significant affinity (as measured using standard radioligand binding assays) or pharmacological activity at 5-HT₂₆, 5-HT₃₇, or 5-HT₄ receptor subtypes or at alpha₂₄, alpha₂₆, or beta-ad-renergic; dopamine₃; dopamine₃; muscarinic; or benzodiaz-emine recentors.

epine receptors. The vascular 5-HT₁ receptor subtype that sumatriptan ac-tivates is present on cranial arteries in both dog and pri-mate, on the human basilar artery, and in the vasculature of human dura mater and mediates vasoconstriction. This action in humans correlates with the relief of migraine head ache. In addition to causing vasoconstriction, experimental data from animal studies show that sumatriptan also activates 5-HT₁ receptors on peripheral terminals of the trigem-inal nerve innervating cranial blood vessels. Such an action may also contribute to the antimigrainous effect of sumatriptan in humans.

In the anesthetized dog, sumatriptan selectively reduces the carotid arterial blood flow with little or no effect on ar-terial blood pressure or total peripheral resistance. In the cat, sumatriptan selectively constrict the carotid arteriove nous anastomoses while having little effect on blood flow or resistance in cerebral or extracerebral tissues.

Pharmacokinetics: The mean maximum concentration following oral dosing with 25 mg is 18 mg/mL (range, 9.747 mg/mL) and 51 ng/mL (range, 9.8400 mg/mL) following oral dosing with 100 mg of summtriptan. This compares with a $C_{\rm max}$ of 5 and 16 ng/mL following dosing with a 5- and 20-mg intranasal dose, respectively. The mean $C_{\rm max}$ following a 6-mg subcutaneous injection is 71 ng/mL (range, 49-110 ng/mL). The bioavailability is approximately 15%, primarily due to presystemic metabolism and partly due to incomplete absorption. The $C_{\rm max}$ is similar during a migraine attack and during a migraine-free period, but the $t_{\rm max}$ is slightly later during the attack, approximately 2.5 hours compared to 2.0 hours. When given as a single dose, sumatriptan display dose proportionality in its extent of Pharmacokinetics: The mean maximum concentration folhours compared to 20 hours. When given as a single cose, sumatriptan display does proportionality in its extent of absorption (area under the curve (AUC)) over the dose range of 25 to 200 mg, but the $C_{\rm max}$ after 100 mg is approx-imately 25% less than expected (based on the 25-mg dose). Food has no significant effect on the bioavailability of sumatriptan, but delays the $t_{\rm max}$ slightly (by about 0.5 hours). hours)

nours). Plasma protein binding is low (14%-21%). The effect of sumatriptan on the protein binding of other drugs has not been evaluated, but would be expected to be minor, given the low rate of protein binding. The apparent volume of dis-

been evaluated, but would be expected to be minor, given the low rate of protein binding. The apparent volume of dis-tribution is 2.4 L/kg. The elimination half-life of sumstriptan is approximately 2.5 hours. Radiolabeled ¹⁴C-sumstriptan administered orally is largely renally excreted (about 60%) with about 40% found in the feces. Most of the radiolabeled compound excreted in the urine is the major metabolite, indole acetic acid (JAA), which is inactive, or the IAA glucuronide. Only 3% of the dose can be recovered as unchanged emitting the second secon

3% of the dose can be recovered as unchanged sumatriptan In vitro studies with human microsomes suggest tha sumatriptan is metabolized by monoamine oxidase (MAO) predominantly the A isoenzyme, and inhibitors of that en-yme may alter sumatriptan pharmacokinetics to increase systemic exposure. No significant effect was seen with an MAO-B inhibitor (see CONTRAINDICATIONS, WARN-

MAN-B INDIDIOT (see CONTRAINDICATIONS, WARN-INGS, and PRECAUTIONS: Drug Interactions). Special Populations: Renal Impairment: The effect of re-nal impairment on the pharmacokinetics of sumatriptan has not been examined, but little clinical effect would be ex-pected as sumatriptan is largely metabolized to an inactive substance.

substance. Mepatic Impairment: The liver plays an important role in the presystemic clearance of orally administered sumatrip-tan. Accordingly, the bioavailability of sumatriptan follow-ing oral administration may be markedly increased in pa-tients with liver disease. In 1 small study of hepatically im-paired patients (n = 8) matched for sex, age, and weight with healthy subjects, the hepatically impaired patients had an approximately 70% increase in AUC and C_{max} and a h_{max} 40 minutes carlier compared to the healthy subjects (see DOSAGE AND ADMINISTRATION).

Age: The pharmacokinetics of oral sumatriptan in the el-derly (mean age, 72 years; 2 males and 4 females) and in patients with migraine (mean age, 35 years; 26 males and 155 females) were similar to that in healthy male subjects (mean age, 30 years) (see PRECAUTIONS: Geriatric Use). Gender: In a study comparing females to males, no phar-maoskinetic differences were observed between genders for AUC, C_{\max} , t_{\max} , and half-life. Alternative structure of sumatriptan Rese: The systemic clearance and C_{\max} of sumatriptan

Rese: The systemic clearance and C_{max} of sumatriptan were similar in black (n = 34) and Caucasian (n = 38) healthy male subjects. Drug intersections: Monoanine Oxidass Inhibitors (MAO9): Treatment with MAO-A inhibitors generally leads to an in-crease of sumatriptan plasma levels (see CONTRAINDICA-TIONS and PRECATIVONS). crease of sumatriptan plasma TIONS and PRECAUTIONS).

Due to gut and hepatic metabolic first-pass effects, the crease of systemic exposure after coadministration of an MAO-A inhibitor with oral sumatriptan is greater than af-ter coadministration of the MAOI with subcutaneous sumatriptan. In a study of 14 healthy females, pretreatment with an MAO-A inhibitor decreased the clearance of subcataneous sumatriptan. Under the conditions of this ex-periment, the result was a 2-fold increase in the area under the sumatriptan plasma concentration x time curve (AUC), corresponding to a 40% increase in elimination half-life. This interaction was not evident with an MAO-B inhibitor. This interaction was not create via a spectra of the second state of the second state

crease in systemic exposure, Alcohol: Alcohol consumed 30 minutes prior to sumatriptan ingestion had no effect on the pharmacokinetics of

CLINICAL STUDIES

The efficacy of IMITREX Tablets in the acute treatment of migraine headaches was demonstrated in 3, randomized double-blind, placebo-controlled studies. Patients enrolled in these 3 studies were predominately female (37%) and Caucasian (97%), with a mean age of 40 years (range, 18-65 years). Patients were instructed to treat a moderate to se-vere 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. Associ-

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Declaration of Rachel J. Watters on Authentication of Publication

1. I, Rachel J. Watters, am a librarian, and the Director of Wisconsin TechSearch ("WTS"), located at 728 State Street, Madison, Wisconsin, 53706. WTS is an interlibrary loan department at the University of Wisconsin-Madison. I have worked as a librarian at the University of Wisconsin library system since 1998. I have been employed at WTS since 2002, first as a librarian and, beginning in 2011, as the Director. Through the course of my employment, I have become well informed about the operations of the University of Wisconsin library system, which follows standard library practices.

2. This Declaration relates to the dates of receipt and availability of the following:

Anonymous. (2010) Zomig Nasal Spray. *Physicians' Desk Reference*, pp. 768-778. Montvale, NJ: PDR Network.

3. <u>Standard operating procedures for materials at the University of</u>

<u>*Wisconsin-Madison Libraries.*</u> When a volume was received by the Library, it would be checked in, stamped with the date of receipt, added to library holdings records, and made available to readers as soon after its arrival as possible. The procedure normally took a few days or at most 2 to 3 weeks.

4. Exhibit A to this Declaration is true and accurate copy of the front matter of the *Physicians' Desk Reference* (2010) publication, which includes a stamp showing that this book is the property of Ebling Library at the University of Wisconsin-Madison.

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Declaration of Rachel J. Watters on Authentication of Publication

Exhibit A also includes an excerpt of pages 768 to 778 of that volume, showing the entry entitled *Zomig Nasal Spray* (2010).

5. Attached as Exhibit B is the cataloging system record of the University of Wisconsin-Madison Libraries for its copy of the *Zomig Nasal Spray* (2010) publication. As shown in the "Receiving date" field of this Exhibit, the University of Wisconsin-Madison Libraries owned this book and had it cataloged in the system as of June 4, 2010.

6. Members of the interested public could locate the *Zomig Nasal Spray* (2010) publication after it was cataloged by searching the public library catalog or requesting a search through WTS. The search could be done by title, author, and/or subject key words. Members of the interested public could access the publication by locating it on the library's shelves or requesting it from WTS.

7. I declare that all statements made herein of my own knowledge are true and that all statements made on information and belief are believed to be true; and further that these statements were made with the knowledge that willful false statements and the like so made are punishable by fine or imprisonment, or both, under Section 1001 of Title 18 of the United States Code.

Date: December 13, 2018 Wisconsin TechSearch

Rachel J. Watter Director

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