HIGHLIGHTS OF PRESCRIBING INFORMATION These highlights do not include all the information

needed to use SUMATRIPTAN INJECTION safely and effectively. See full prescribing information for SUMATRIPTAN INJECTION. SUMATRIPTAN injection, for subcutaneous use

Initial U.S. Approval: 1992 ----RECENT MAJOR CHANGES -Dosage and Administration Dosing Information (2.1) Removal of Dosage and Administration,

 $Administration \, of \, Doses \, of \, sum a tript an \, injection$ Other than 6 mg (2.3) 12/2021 -----INDICATIONS AND USAGE--Sumatriptan injection is a serotonin (5-HT_{1B/ID}) receptor agonist (triptan) indicated for:

 Acute treatment of migraine with or without aura in adults (1) Acute treatment of cluster headache in adults (1)

Limitations of Use: Use only if a clear diagnosis of migraine or

 Not indicated for the prophylactic therapy of migraine or cluster headache attacks (1) ---- DOSAGE AND ADMINISTRATION --For subcutaneous use only (2.1)

cluster headache has been established (1)

 Acute treatment of migraine: single dose of 1 to 6 mg (2.1) Acute treatment of cluster headache: single

dose of 6 mg (2.1) separate doses by at least 1 hour (2.1) -----DOSAGE FORMS AND STRENGTHS-----

 Injection: 6 mg single-dose prefilled syringe assembled in an autoinjector (3) ----CONTRAINDICATIONS-- History of coronary artery disease or coronary artery vasospasm (4)

 Wolff-Parkinson-White syndrome or other cardiac accessory conduction pathway disorders (4) History of stroke, transient ischemic attack, or

hemiplegic or basilar migraine (4) Peripheral vascular disease (4) Ischemic bowel disease (4) Uncontrolled hypertension (4)

FULL PRESCRIBING INFORMATION: CONTENTS* 1 INDICATIONS AND USAGE

2.1 Dosing Information 2.2 Administration Using the Sumatriptan 8 Autoinjector

4 CONTRAINDICATIONS 5 WARNINGS AND PRECAUTIONS 5.1 Myocardial Ischemia, Myocardia Infarction, and Prinzmetal's Angina

3 DOSAGE FORMS AND STRENGTHS

2 DOSAGE AND ADMINISTRATION

5.2 Arrhythmias 5.3 Chest, Throat, Neck and/or Jaw Pain/Tightness/Pressure 5.4 Cerebrovascular Events 5.5 Other Vasospasm Reactions

5.7 Serotonin Syndrome 5.8 Increase in Blood Pressure 5.9 Hypersensitivity Reactions 5.10 Seizures 6 ADVERSE REACTIONS

administered to treat any subsequent attacks.

2.2 Administration Using the Sumatriptan Autoinjector

stroke [see Warnings and Precautions (5.4)].

Peripheral vascular disease [see Warnings and Precautions (5.5)] • Ischemic bowel disease [see Warnings and Precautions (5.5)].

• Sumatriptan injection is not indicated for the prevention of migraine or cluster headache attacks

injection sites with an adequate skin and subcutaneous thickness to accommodate the length of the needle.

 $For the \, treatment \, of \, cluster \, headache, the \, efficacy \, of \, lower \, doses \, has \, not \, been \, established.$

5.6 Medication Overuse Headache

6.1 Clinical Trials Experience 6.2 Postmarketing Experience 7 DRUGINTERACTIONS 7.1 Ergot-Containing Drugs

FULL PRESCRIBING INFORMATION

2 DOSAGE AND ADMINISTRATION 2.1 Dosing Information

3 DOSAGE FORMS AND STRENGTHS

4 CONTRAINDICATIONS

Precautions (5.2)].

1 INDICATIONS AND USAGE

headache. Limitations of Use: Recent (within 24 hours) use of another 5-HT. agonist (e.g., another triptan) or of an ergotaminecontaining medication (4) Concurrent or recent (past 2 weeks) use of

monoamine oxidase-Ainhibitor (4) Hypersensitivity to sumatriptan injection (angioede Severe hepatic impairment [see Clinical Pharmacology (12.3)]. Hypersensitivity to sumatriptan injection (angioedema and anaphylaxis seen) (4) 5 WARNINGS AND PRECAUTIONS Severe hepatic impairment (4) 5.1 Myocardial Ischemia, Myocardial Infarction, and Prinzmetal's Angina

The use of sumatriptan injection is contraindicated in patients with ischemic or vasospastic CAD. There have been rare reports of serious cardia ----- WARNINGS AND PRECAUTIONS -----adverse reactions, including acute myocardial infarction, occurring within a few hours following administration of sumatriptan injection. Some of Myocardial ischemia/infarction and Prinzmetal's angina: Perform cardiac evaluation in patients patients without a history of CAD. with multiple cardiovascular risk factors (5.1) Perform a cardiovascular evaluation in triptan-naive patients who have multiple cardiovascular risk factors (e.g., increased age, diabetes, hypertension, smoking, obesity, strong family history of CAD) prior to receiving sumatriptan injection. If there is evidence of CAD or coronary artery Arrhythmias: Discontinue sumatriptan injection vasospasm, sumatriptan injection is contraindicated. For patients with multiple cardiovascular risk factors who have a negative cardiov

 Chest/throat/neck/jaw pain, tightness, pressure, or heaviness: Generally not associated long-term users of sumatriptan injection. with myocardial ischemia; evaluate for coronary artery disease in patients at high risk (5.3) Life-threatening disturbances of cardiac rhythm, including ventricular tachycardia and ventricular fibrillation leading to death, have been reported

if occurs (5.2)

Cerebral hemorrhage, subarachnoid hemorrhage, and stroke: Discontinue sumatriptan injection if Gastrointestinal ischemic reactions and peripheral vasospastic reactions: Discontinue

sumatriptan injection if occurs (5.5) Medication overuse headache: Detoxification 5.4 Cerebrovascular Events Cerebral hemorrhage, subarachnoid hemorrhage, and stroke have occurred in patients treated with 5-HT, agonists, and some have resulted in may be necessary (5.6) fatalities. In a number of cases, it appears possible that the cerebrovascular events were primary, the 5-HT, agonist having been administered in the incorrect belief that the symptoms experienced were a consequence of migraine when they were not. Also, patients with migraine may be at Serotonin syndrome: Discontinue sumatriptan injection if occurs (5.7) icreased risk of certain cerebrovascular events (e.g., stroke, hemorrhage, TIA). Discontinue sumatriptan injection if a cere

epilepsy or a lowered seizure threshold (5.10) ----- ADVERSE REACTIONS --- Maximum dose in a 24-hour period: 12 mg, Most common adverse reactions (≥ 5% and 5.5 Other Vasospasm Reactions Sumatriptan injection may cause non-coronary vasospastic reactions, such as peripheral vascular ischemia, gastrointestinal vascular ischemia and > placebo) were injection site reactions, tingling, infarction (presenting with abdominal pain and bloody diarrhea), splenic infarction, and Raynaud's syndrome. In patients who experience symptoms or signs suggestive of non-coronary vasospasm reaction following the use of any 5-HT1 agonist, rule out a vasospastic reaction before receiving dizziness/vertigo, warm/hot sensation, burning sensation, feeling of heaviness, pressure sensation, Reports of transient and permanent blindness and significant partial vision loss have been reported with the use of 5-HT, agonists. Since visual flushing, feeling of tightness, and numbness (6.1) To report SUSPECTED ADVERSE REACTIONS, 5.6 Medication Overuse Headache

Seizures: Use with caution in patients with

contact Dr. Reddy's Laboratories Inc. at 1-888-375-3784 or FDA at 1-800-FDA-1088 or www.fda.gov/medwatch. ---- USE IN SPECIFIC POPULATIONS ----Pregnancy: Based on animal data, may cause fetal

See 17 for PATIENT COUNSELING INFORMATION and FDA- approved patient labeling.

and Serotonin Syndrome

USE IN SPECIFIC POPULATIONS

8.1 Pregnancy

8.4 Pediatric Use

8.5 Geriatric Use

12 CLINICAL PHARMACOLOGY

12.3 Pharmacokinetics

13 NONCLINICAL TOXICOLOGY

14 CLINICAL STUDIES

14.2 Cluster Headache

17 PATIENT COUNSELING INFORMATION

prescribing information are not listed.

14.1 Migraine

Sum a tript an injection is indicated in a dults for (1) the acute treatment of migraine, with or without aura, and (2) the acute treatment of cluster and (3) the acute treatment of cluster and (4) the acute treatment of cluster and (5) the acute treatment of cluster and (6) the acute treatment of cluster and (7) the acute treatment of cluster and (8) the acute tre

Use only if a clear diagnosis of migraine or cluster headache has been established. If a patient has no response to the first

The maximum single recommended adult dose of sumatriptan injection for the acute treatment of migraine or cluster headache is 6 mg injected subcutaneously. For the treatment of migraine, if side effects are dose limiting, the lower dose of 4 mg may be used [see Clinical Studies (14.1)].

The maximum cumulative dose that may be given in 24 hours is 12 mg, two 6 mg injections separated by at least 1 hour. A second 6 mg dose should

An autoinjector device (sumatriptan autoinjector) is available for use with 6 mg prefilled syringe to facilitate self-administration in patients using the 6 mg dose. With this device, the needle penetrates approximately 1/4 inch (5 to 6 mm). The injection is intended to be given subcutaneously, and

intramuscular or intravascular delivery must be avoided. Instruct patients on the proper use of sumatriptan autoinjector and direct them to use

Injection: 6 mg prefilled syringe assembled in an Autoinjector. Each 0.5 mL injection contains 8.4 mg of sumatriptan succinate, USP equivalent
to 6 mg of sumatriptan.

Ischemic coronary artery disease (CAD) (angina pectoris, history of myocardial infarction, or documented silent ischemia) or coronary artery
vasospasm, including Prinzmetal's angina [see Warnings and Precautions (5.1)]

History of stroke or transient ischemic attack (TIA) or history of hemiplegic or basilar migraine because these patients are at a higher risk of

migraine or cluster headache attack treated with sumatriptan injection, reconsider the diagnosis before sumatriptan injection is

12.1 Mechanism of Action

12.2 Pharmacodynamics

8.2 Lactation

OVERDOSAGE

11 DESCRIPTION

7.2 Monoamine Oxidase-Alnhibitors 7.3 Other 5-HT, Agonists 7.4 Selective Serotonin Reuptake Inhibitors/ Serotonin Norepinephrine Reuptake Inhibitors

Revised: 12/2022

life-threatening or fatal. In general, anaphylactic reactions to drugs are more likely to occur in individuals with a history of sensitivity to multiple

Recent use (i.e., within 24 hours) of ergotamine-containing medication, ergot-type medication (such as dihydroergotamine or methysergide), or another 5-hydroxytryptamine, (5-HT,) agonist [see Drug Interactions (7.1, 7.3)].

Interactions (7.2), Clinical Pharmacology (12.3)].

5.3 Chest, Throat, Neck, and/or Jaw Pain/Tightness/Pressure

additional injections of sumatriptan.

5.8 Increase in Blood Pressure

5.9 Hypersensitivity Reactions

Concurrent administration of a monoamine oxidase (MAO)-A inhibitor or recent (within 2 weeks) use of an MAO-A inhibitor [see Drug

evaluation, consider administering the first dose of sumatriptan injection in a medically supervised setting and performing an electrocardiogram

within a few hours following the administration of 5-HT, agonists. Discontinue sumatriptan injection if these disturbances occur. Sumatriptan

injection and are usually non-cardiac in origin. However, perform a cardiac evaluation if these patients are at high cardiac risk. The use of

Before treating headaches in patients not previously diagnosed with migraine or cluster headache or in patients who present with atypical

symptoms, exclude other potentially serious neurological conditions. Sumatriptan injection is contraindicated in patients with a history of stroke or

disorders may be part of a migraine attack, a causal relationship between these events and the use of 5-HT, agonists has not been clearly

Overuse of acute migraine drugs (e.g., ergotamine, triptans, opioids, or combination of these drugs for 10 or more days per month) may lead to exacerbation of headache (medication overuse headache). Medication overuse headache may present as migraine-like daily headaches or as a marked increase in frequency of migraine attacks. Detoxification of patients, including withdrawal of the overused drugs, and treatment of

serotonin norepinephrine reuptake inhibitors (SNRIs), tricyclic antidepressants (TCAs), and MAO inhibitors [see Drug Interactions (7.4)]. Serotonin pyndrome symptoms may include mental status changes (e.g., agitation, hallucinations, coma), autonomic instability (e.g., tachycardia, labile blood pressure, hyperthermia), neuromuscular aberrations (e.g. hyperreflexia, incoordination), and/or gastrointestinal symptoms (e.g., nausea, vomiting,

iarrhea). The onset of symptoms usually occurs within minutes to hours of receiving a new or a greater dose of a serotonergic medication.

Significant elevation in blood pressure, including hypertensive crisis with acute impairment of organ systems, has been reported on rare occasio

in patients treated with 5-HT, agonists, including patients without a history of hypertension. Monitor blood pressure in patients treated with sumatriptan injection. Sumatriptan injection is contraindicated in patients with uncontrolled hypertension.

injection is contraindicated in patients with Wolff-Parkinson-White syndrome or arrhythmias associated with other cardiac accessory conduction

Sensations of tightness, pain, pressure, and heaviness in the precordium, throat, neck, and jaw commonly occur after treat

Seizures have been reported following administration of sumatriptan injection. Some have occurred in patients with either a history of seizures or concurrent conditions predisposing to seizures. There are also reports in patients where no such predisposing factors are apparent. Sumatriptan injection should be used with caution in patients with a history of epilepsy or conditions associated with a lowered seizure threshold. 6 ADVERSEREACTIONS

The following serious adverse reactions are described below and elsewhere in the labeling: Myocardial ischemia, myocardial infarction, and Prinzmetal's angina [see Warnings and

Arrhythmias [see Warnings and Precautions (5.2)] Chest, throat, neck, and/or jaw pain/tightness/pressure [see Warnings and Precautions (5.3)] Cerebrovascular events [see Warnings and Precautions (5.4)] Other vasos pasm reactions [see Warnings and Precautions (5.5)]

Medication overuse headache [see Warnings and Precautions (5.6)] Serotonin syndrome [see Warnings and Precautions (5.7)] Increase in blood pressure [see Warnings and Precautions (5.8)] Hypersensitivity reactions [see Contraindications (4), Warnings and Precautions (5.9)]

13.1 Carcinogenesis, Mutagenesis, Seizures [see Warnings and Precautions (5.10)] Impairment of Fertility 6.1 Clinical Trials Experience 13.2 Animal Toxicology and/or Pharmacology Because clinical trials are conducted under widely varying conditions, adverse reaction rates observed in the clinical trials of a drug cannot be

directly compared with rates in the clinical trials of another drug and may not reflect the rates observed in practice Table 1 lists adverse reactions that occurred in 2 U.S. placebo-controlled clinical trials in patients with migraines (Studies 2 and 3) following either a 16 HOW SUPPLIED/STORAGE AND HANDLING

 $single\ 6\ mg\ dose\ of\ sum a triptan\ injection\ or\ placebo.\ Only\ reactions\ that\ occurred\ at\ a\ frequency\ of\ 2\%\ or\ more\ in\ groups\ treated\ with\ sum a triptan\ of\ 2\%\ or\ more\ in\ groups\ treated\ with\ sum a triptan\ of\ 2\%\ or\ more\ in\ groups\ treated\ with\ sum a triptan\ of\ 2\%\ or\ more\ in\ groups\ treated\ with\ sum a triptan\ of\ 2\%\ or\ more\ in\ groups\ treated\ with\ sum a triptan\ of\ 2\%\ or\ more\ in\ groups\ treated\ with\ sum a triptan\ of\ 2\%\ or\ more\ in\ groups\ treated\ of\ 2\%\ or\ more\ in\ groups\ treated\ of\ 2\%\ or\ more\ of\ 2\%\ or\ of\ 2\%\ of\ 2\%\ or\ of\ 2\%\ of\$ jection 6 mg and that occurred at a frequency greater than the placebo group are included in Table 1. Table 1. Adverse Reactions in Pooled Placebo-Controlled Trials in Patients with Migraine (Studies 2 and 3) *Sections or subsections omitted from the full Sumatriptan Injection

	6 mg Subcutaneous	(n = 370)
Adverse Reaction	(n = 547)	(11 - 370)
narono noaction	%	,,,
Atypical sensations	42	9
Tingling	14	3
Warm/hot sensation	11	4
Burning sensation	7	<1
Feeling of heaviness	7	1
Pressure sensation	7	2
Feeling of tightness	5	<1
Numbness	5	2
Feeling strange	2	<1
Tight feeling in head	2	<1
Cardiovascular	-	
Flushing	7	2
Chest discomfort	5	1 1
Tightness in chest	3	4
Pressure in chest	2	<1
Ear, nose, and throat Throat discomfort	_	
	3	4
Discomfort: nasal cavity/sinuses	2	<1
Injection site reaction ^a	59	24
Miscellaneous		
Jaw discomfort	2	0
Musculoskeletal		
Weakness		
Neck pain/stiffness	5	<1
·	5	<1
Myalgia	2	<1
Neurological		
Dizziness/vertigo	12	4
Drowsiness/sedation	3	2
Headache	2	<1
Skin		
Sweating	2	1

 $Includes\,injection\,site\,pain, stinging/burning, swelling, erythema, bruising, bleeding$ The incidence of adverse reactions in controlled clinical trials was not affected by gender or age of the patients. There were insufficient data to

In the controlled clinical trials assessing the efficacy of sumatriptan injection as a treatment for cluster headache (Studies 4 and 5), no new Overall, the frequency of adverse reactions reported in the trials of cluster headache was generally lower than in the migraine trials. Exception include reports of paresthesia (5% sumatriptan injection, 0% placebo), nausea and vomiting (4% sumatriptan injection, 0% placebo), and

6.2 Postmarketing Experience The following adverse reactions have been identified during postapproval use of sumatriptan tablets, sumatriptan nasal spray, and sumatriptan injection. Because these reactions are reported voluntarily from a population of uncertain size, it is not always possible to reliably estimate their

Hypotension, palpitations

ontaining or ergot-type medications (like dihydroergotamine or methysergide) and sumatriptan injection within 24 hours of each other is

MAO-A inhibitors increase systemic exposure by 2-fold. Therefore, the use of sumatriptan injection in patients receiving MAO-A inhibitors is contraindicated [see Clinical Pharmacology (12.3)]. 7.3 Other 5-HT, Agonists

Because their vasospastic effects may be additive, coadministration of sumatriptan injection and other 5-HT, agonists (e.g., triptans) within 7.4 Selective Serotonin Reuptake Inhibitors/Serotonin Norepinephrine Reuptake Inhibitors and Serotonin Syndrome
Cases of serotonin syndrome have been reported during coadministration of triptans and SSRIs, or SNRIs, TCAs, and MAO inhibitors [see Warnings

8 USE IN SPECIFIC POPULATIONS

Data from a prospective pregnancy exposure registry and epidemiological studies of pregnant women have not detected an increased frequency of birth defects or a consistent pattern of birth defects among women exposed to sumatriptan compared with the general population (see Data). In developmental toxicity studies in rats and rabbits, oral administration of sumatriptan to pregnant animals was associated with embryolethality, In the U.S. general population, the estimated background risk of major birth defects and of miscarriage in clinically recognized pregnancies is 2% to 4% and 15% to 20%, respectively. The reported rate of major birth defects among deliveries to women with migraine ranged from 2.2% to 2.9% and the reported rate of miscarriage was 17%, which were similar to rates reported in women without migraine. Disease-Associated Maternal and/or Embryo/Fetal Risk: Several studies have suggested that women with migraine may be at increased risk of

Human Data: The Sumatriptan/Naratriptan/TREXIMET (sumatriptan and naproxen sodium) Pregnancy Registry, a population-based international prospective study, collected data for sumatriptan from January 1996 to September 2012. The Registry documented outcomes of 626 infants and fetuses exposed to sumatriptan during pregnancy (528 with earliest exposure during the first trimester, 78 during the second trimester, 16 during the third trimester, and 4 unknown). The occurrence of major birth defects (excluding fetal deaths and induced abortions without reported defects (excluding fetal deaths and induced abortions without reported defects (excluding fetal deaths and induced abortions without reported defects (excluding fetal deaths and induced abortions without reported defects (excluding fetal deaths and induced abortions without reported defects (excluding fetal deaths and induced abortions without reported defects (excluding fetal deaths and induced abortions without reported defects (excluding fetal deaths and induced abortions without reported defects (excluding fetal deaths and induced abortions without reported defects (excluding fetal deaths and induced abortions without reported defects (excluding fetal deaths and induced abortions without reported defects (excluding fetal deaths are fetal deaths and induced abortions without reported defects (excluding fetal deaths are fand all spontaneous pregnancy losses) during first-trimester exposure to sumatriptan was 4.2% (20/478 [95% Cl: 2.6% to 6.5%]) and during any trimester of exposure was 4.2% (24/576 [95% Cl: 2.7% to 6.2%]). The sample size in this study had 80% power to detect at least a 1.73-to 1.91-fold increase in the rate of major malformations. The number of exposed pregnancy outcomes accumulated during the registry was insufficient to support definitive conclusions about overall malformation risk or for making comparisons of the frequencies of specific birth defects. Of the 20 infants with reported birth defects after exposure to sumatriptan in the first trimester, 4 infants had ventricular septal defects, including one infant who was exposed to both sumatriptan and naratriptan, and 3 infants had pyloric stenosis. No other birth defect was reported for more than 2 infants

In a study using data from the Swedish Medical Birth Register, live births to women who reported using triptans or ergots during pregnancy were compared with those of women who did not. Of the 2,257 births with first-trimester exposure to sumatriptan, 107 infants were born with malformations (relative risk 0.99 [95% Cl: 0.91 to 1.21]). A study using linked data from the Medical Birth Registry of Norway to the Norwegian Prescription Database compared pregnancy outcomes in women who redeemed prescriptions for triptans during pregnancy, as well as a migraine disease comparison group who redeemed prescriptions for sumatriptan before pregnancy only, compared with a population control group. Of the escriptions for sum a triptan during the first trimester, 15 had infants with major congenital malformations (OR 1.16 [95% and 1.16 [95% are not only the first trimester). The sum of the first trimester is a sum of the first trimester in the first trCl: 0.69 to 1.94]) while for the 364 women who redeemed prescriptions for sumatriptan before, but not during, pregnancy, 20 had infants with major nital malformations (OR 1.83 [95% CI: 1.17 to 2.88]), each compared with the population comparison group. Additional smaller observational studies evaluating use of sumatriptan during pregnancy have not suggested an increased risk of teratogenicity.

Animal Data: Oral administration of sumatriptan to pregnant rats during the period of organogenesis resulted in an increased incidence of fetal blood vessel (cervicothoracic and umbilical) abnormalities. The highest no-effect dose for embryofetal developmental toxicity in rats was 60 mg/kg/day. Oral administration of sumatriptan to pregnant rabbits during the period of organogenesis resulted in increased incidences of embryolethality and fetal cervicothoracic vascular and skeletal abnormalities. Intravenous administration of sumatriptan to pregnant rabbits during the period of organogenesis resulted in an increased incidence of embryolethality. The highest oral and intravenous no-effect doses for developbits were 15 and 0.75 mg/kg/day, respectively.

Oral administration of sumatriptan to rats prior to and throughout gestation resulted in embryofetal toxicity (decreased body weight, decreased ossification, increased incidence of skeletal abnormalities). The highest no-effect dose was 50 mg/kg/day. In offspring of pregnant rats treated orally with sumatriptan during organogenesis, there was a decrease in pup survival. The highest no-effect dose for this effect was 60 mg/kg/day. The highest no-effect dose for this finding was 100 mg/kg/day 8.2 Lactation

Risk Summary Sumatriptan is excreted in human milk following subcutaneous administration (see Data). There are no data on the effects of sumatriptan on the The developmental and health benefits of breastfeeding should be considered along with the mother's clinical need for sumatriptan injection and any potential adverse effects on the breastfed infant from sumatriptan or from the underlying maternal condition

Clinical Considerations

8.4 Pediatric Use Safety and effectiveness in pediatric patients have not been established. Sumatriptan injection is not recommended for use in patients younge than 18 years of age. Two controlled clinical trials evaluated sumatriptan nasal spray (5 to 20 mg) in 1,248 pediatric migraineurs aged 12 to 17 years who treated a single attack. The trials did not establish the efficacy of sumatriptan nasal spray compared with placebo in the treatment of migraine in pediatric satients. Adverse reactions observed in these clinical trials were similar in nature to those reported in clinical trials in adults

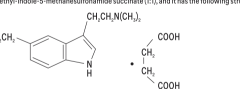
Five controlled clinical trials (2 single-attack trials, 3 multiple-attack trials) evaluating gral sumatriptan (25 to 100 mg) in pediatric patients aged 12 to 17 years enrolled a total of 701 pediatric migraineurs. These trials did not establish the efficacy of oral sumatriptan compared with placebo in the treatment of migraine in pediatric patients. Adverse reactions observed in these clinical trials were similar in nature to those reported in clinical trials in adults. The frequency of all adverse reactions in these patients appeared to be both dose- and age-dependent, with younger patients reporting reactions more commonly than older pediatric patients. Postmarketing experience documents that serious adverse reactions have occurred in the pediatric population after use of subcutaneous, oral, and/or intranasal sumatriptan. These reports include reactions similar in nature to those reported rarely in adults, including stroke, visual loss, and death. A myocardial infarction has been reported in a 14-year-old male following the use of oral sumatriptan; clinical signs occurred within 1 day of

8.5 Geriatric Use Clinical trials of sumatriptan injection did not include sufficient numbers of patients aged 65 years and older to determine whether they respond differently from younger patients. Other reported clinical experience has not identified differences in responses between the elderly and younger patients. In general, dose selection for an elderly patient should be cautious, usually starting at the low end of the dosing range, reflecting the greater frequency of decreased hepatic, renal, or cardiac function and of concomitant disease or other drug therapy. A cardiov a scular evaluation is recommended for geriatric patients who have other cardiov a scular risk factors (e.g., diabetes, hypertension, smoking, and the scular risk factors (e.g., diabetes, hypertension, smoking, factors (e.g., diabetes, hypertension), smoking, and the scular risk factors (e.g., diabetes, hypertension), smoking, and the scular risk factors (e.g., diabetes, hypertension), smoking, and the scular risk factors (e.g., diabetes, hypertension), smoking, and the scular risk factors (e.g., diabetes, hypertension), smoking, and the scular risk factors (e.g., diabetes, hypertension), smoking, and the scular risk factors (e.g., diabetes, hypertension), smoking, and the scular risk factors (e.g., diabetes, hypertension), smoking, and the scular risk factors (e.g., diabetes, hypertension), smoking, and the scular risk factors (e.g., diabetes, hypertension), and the scular risk faobesity, strong family history of CAD) prior to receiving sumatriptan injection [see Warnings and Precautions (5.1)].

drug administration. Clinical data to determine the frequency of serious adverse reactions in pediatric patients who might receive subcutaneous,

Coronary vasospasm was observed after intravenous administration of sumatriptan injection [see Contraindications (4)]. Overdoses would be expected from animal data (dogs at 0.1 g/kg, rats at 2 g/kg) to possibly cause convulsions, tremor, inactivity, erythema of the extremities, reduced espiratory rate, cyanosis, ataxia, mydriasis, injection site reactions (desquamation, hair loss, and scab formation), and paralysis The elimination half-life of sumatriptan is about 2 hours [see Clinical Pharmacology (12.3)]; therefore, monitoring of patients after overdose with sumatriptan injection should continue for at least 10 hours or while symptoms or signs persist. t is unknown what effect hemodialysis or peritoneal dialysis has on the serum concentrations of sumatrip

Sumatriptan injection, USP contains sumatriptan succinate, USP a selective $5\text{-HT}_{\text{luco}}$ receptor agonist. Sumatriptan succinate, USP is chemically designated as 3-[2-(dimethylamino)ethyl]-N- methyl-indole-5- methanesulfonamide succinate (1:1), and it has the following structure: CH, NHSO, CH,



The molecular formula is C., H., N.O., S. • C. H.O., representing a molecular weight of 413.5. Sumatriptan succinate, USP is a white or almost white Sumatriptan injection, USP is a clear, colorless to pale yellow, free from visible particulate matter, sterile nonpyrogenic solution for subcutaneo

Sumatriptan binds with high affinity to human cloned 5-HT_{18/10} receptors. Sumatriptan presumably exerts its therapeutic effects in the treatr migraine and cluster headaches through agonist effects at the 5-HT_{IB/ID} receptors on intracranial blood vessels and sensory nerves of the trigeminal system, which result in cranial vessel constriction and inhibition of pro-inflammatory neuropeptide release

Significant elevation in blood pressure, including hypertensive crisis, has been reported in patients with and without a history of hypertension [see Warnings and Precautions (5.8)].

Peripheral (Small) Arteries In healthy volunteers (N = 18), a trial evaluating the effects of sumatriptan on peripheral (small vessel) arterial reactivity failed to detect a clinically

Heart Rate Transient increases in blood pressure observed in some patients in clinical trials carried out during sumatriptan's development as a treatment for migraine were not accompanied by any clinically significant changes in heart rate.

12.3 Pharmacokinetics The bioavailability of sumatriptan via subcutaneous site injection to 18 healthy male subjects was 97% ± 16% of that obtained following intravenous After a single 6 mg subcutaneous manual injection into the deltoid area of the arm in 18 healthy males (age: 24 ± 6 years, weight: 70 kg), the

maximum serum concentration (C_{max}) of sumatriptan was (mean \pm standard deviation) 74 \pm 15 ng/mL and the time to peak concentration (T_{max}) was 12 minutes after injection (range: 5 to 20 minutes). In this trial, the same dose injected subcutaneously in the thigh gave a C_{max} of 61 \pm 15 ng/mL by manual injection versus 52 \pm 15 ng/mL by autoinjector techniques. The T_{max} or amount absorbed was not significantly altered by either the site or $Protein \ binding, determined \ by \ equilibrium \ dialysis \ over the \ concentration \ range \ of 10 \ to 1,000 \ ng/mL \ is \ low, approximately 14\% \ to 21\%. \ The \ effect \ of \ binding \ approximately 14\% \ to 21\%.$ umatriptan on the protein binding of other drugs has not been evaluated.

Following a 6 mg subcutaneous injection into the deltoid area of the arm in 9 males (mean age: 33 years, mean weight: 77 kg) the volume of $In\ vitro\ studies\ with\ human\ microsomes\ suggest\ that\ sumatriptan\ is\ metabolized\ by\ MAO,\ predominantly\ the\ A\ isoenzyme.\ Most\ of\ a\ radiolabeled\ dose\ of\ sumatriptan\ excreted\ in\ the\ urine\ is\ the\ major\ metabolite\ indole\ acetic\ acid\ (IAA)\ or\ the\ IAA\ glucuronide,\ both\ of\ which\ are\ in\ active.$

After a single 6 mg subcutaneous dose, 22% ± 4% was excreted in the urine as unchanged sumatriptan and 38% ± 7% as the IAA metabolite Following a 6 mg subcutaneous injection into the deltoid area of the arm, the systemic clearance of sumatriptan was 1,194 ± 149 mL/min and the terminal half-life was 115 ± 19 minutes.

Specific Populations Age: The pharmacokinetics of sumatriptan in the elderly (mean age: 72 years, 2 males and 4 females) and in subjects with migraine (mean age: 38 years, 25 males and 155 females) were similar to that in healthy male subjects (mean age: 30 years).

Patients with Hepatic Impairment: The effect of mild to moderate hepatic disease on the pharmacokinetics of subcutaneously administered sumatriptan has been evaluated. There were no significant differences in the pharmacokinetics of subcutaneously administered sumatriptan in moderately hepatically impaired subjects compared with healthy controls. The pharmacokinetics of subcutaneously administered sumatriptan in patients with severe hepatic impairment has not been studied. The use of sumatriptan injection in this population is contraindicated [see $\textit{Racial Groups:} \ \text{The systemic clearance and C_{\max} of subcutaneous sumatriptan were similar in black (n = 34) and $Caucasian (n = 38)$ healthy male}$

Monoamine Oxidase-A Inhibitors: In a trial of 14 healthy females, pretreatment with an MAO-A inhibitor decreased the clearance of subcutaneous sumatriptan, resulting in a 2-fold increase in the area under the sumatriptan plasma concentration-time curve (AUC), corresponding to a 40% increase in elimination half-life. 13 NONCLINICAL TOXICOLOGY

13.1 Carcinogenesis, Mutagenesis, Impairment of Fertility

160 mg/kg/day (the high dose in rat was reduced from 360 mg/kg/day during Week 21). The highest dose to mice and rats was approximately 130 and 260 times the single MRHD of 6 mg administered subcutaneously on a mg/m² basis. There was no evidence in either species of an increase in tumors related to sumatriptan administration

Sumatriptan was negative in in vitro (bacterial reverse mutation [Ames], gene cell mutation in Chinese hamster V79/HGPRT, chromosomal Impairment of Fertility

treatment-related decrease in fertility secondary to a decrease in mating in animals treated with doses greater than 5 mg/kg/day. It is not clear whether this finding was due to an effect on males or females or both. When sumatriptan was administered by subcutaneous injection to male and female rats prior to and throughout the mating period, there was no vidence of impaired fertility at doses up to 60 mg/kg/day.

ogs receiving oral sumatriptan developed corneal opacities and defects in the corneal epithelium. Corneal opacities were seen at the lowest dose tested, 2 mg/kg/day, and were present after 1 month of treatment. 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 plasma exposure at the

14 CLINICAL STUDIES 14.1 Migraine

of the symptoms enumerated in Table 3, onset of relief began as early as 10 minutes following a 6 mg sumatriptan injection. Lower doses of sumatriptan injection may also prove effective, although the proportion of patients obtaining adequate relief was decreased and the latency to that In Study 1, 6 different doses of sumatriptan injection (n = 30 each group) were compared with placebo (n = 62), in a single-attack, parallel-group design, the dose-response relationship was found to be as shown in Table 2. 2. Proportion of Patients with Migraine Relief and Incidence of Adverse Reactions by Time and by Sumatriptan Dose in Study 1

Table 2. Proportion of Patients with Migraine Relief and Incidence of Adverse Reactions by Time and by Sumatriptan Dose in Study 1											
Dose of Sumatriptan Injection			Percent Patients with R	nts with Relief" Adverse Reactions							
,	at 10 Minutes	at 30 Minutes	at 1 Hour	at 2 Hours	Incidence (%)						
Placebo	5	15	24	21	55						
1 mg	10	40	43	40	63						
2 mg	7	23	57	43	63						
3 mg	17	47	57	60	77						
4 mg	13	37	50	57	80						
6 mg	10	63	73	70	83						
8 mg	23	57	80	83	93						
*Poliofic defined as the red	uction of moderate or co	wara pain to no ar mild	nain aftar daoing witho	utuse of resous medica	tion						

In 2 randomized, placebo-controlled clinical trials of sumatriptan injection 6 mg in 1,104 patients with moderate or severe migraine pain (Studies 2 and 3), the onset of relief was less than 10 minutes. Headache relief, as defined by a reduction in pain from severe or moderately severe to mild or no headache, was achieved in 70% of the patients within 1 hour of a single 6 mg subcutaneous dose of sumatriptan injection. Approximately 82% and 65% of patients treated with sumatriptan injection 6 mg had headache relief and were pain free within 2 hours, respectively.

1 -Hour Data		Study 2	Study 3			
	Placebo (n = 190)	Sumatriptan Injection 6 mg (n = 384)	Placebo (n = 180)	Sumatriptan Injection 6 mg (n = 350)		
Patients with pain relief (Grade 0/1)	18%	70% ^a	26%	70%ª		
Patients with no pain	5%	48% a	13%	49% a		
Patients without nausea	48%	73% ^a	50%	73% ^a		
Patients without photophobia	23%	56% a	25%	58% a		
Patients with little or no clinical disability ^b	34%	76% a	34%	76% a		
2-Hour Data	:	Study 2	Study 3			
	Placebo ^c	Sumatriptan Injection 6 mg ^d	Placebo ^c	Sumatriptan Injection 6 mg ^d		
Patientswith pain relief (Grade 0/1)	31%	81%ª	39%	82%ª		
Patients with no pain	11%	63%ª	19%	65% a		
Patients without nausea	56%	82% ^a	63%	81% a		
Patients without photophobia	31%	72% ^a	35%	71% a		
Patients with little or no clinical disability ^b	42%	85% a	49%	84% a		

Basuccessful outcome in terms of clinical disability was defined prospectively as ability to work mildly impaired or ability to work and function °Includes patients that may have received an additional placebo injection 1 hour after the initial injection

⁴Includes patients that may have received an additional 6 mg of sumatriptan injection 1 hour after the initial injectio

reactions.				injection. Each 0.5 mL of sumatriptan 3.5 mg of sodium chloride, USP in Wat between 275 and 315 m0sm/kg.										
				PHARMACIST - DETACH FROM HERE										
too hes 'se, ent	ous tan nti-	the	ere	t ing of of	hat	on.	ე"(ase	Jot	of	nan nan not not not not not not not not not no	ers The rse	pur	

10 OVERDOSAGE

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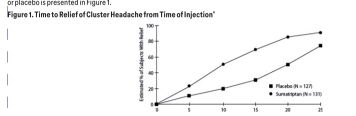
umatriptan injection also relieved photophobia, phonophobia (sound sensitivity), nausea, and vomiting associated with migraine attacks. The efficacy of sumatriptan injection was unaffected by whether or not the migraine was associated with aura, duration of attack, gender or age of

an allergy to sumatriptan or any of the ingredients in sumatriptan injection. See the end of this leaflet for a complete list of ingredients in the patient, or concomitant use of common migraine prophylactic drugs (e.g., beta-blockers).

14.2 Cluster Headache The efficacy of sumatriptan injection in the acute treatment of cluster headache was demonstrated in 2 randomized, double-blind, placebocontrolled, 2-period crossover trials (Studies 4 and 5). Patients aged 21 to 65 years were enrolled and were instructed to treat a moderate to very severe headache within 10 minutes of onset. Headache relief was defined as a reduction in headache severity to mild or no pain. In both trials, the proportion of individuals gaining relief at 10 or 15 minutes was significantly greater among patients receiving 6 mg of sumatriptan injection npared with those who received placebo (see Table 4). Table 4. Proportion of Patients with Cluster Headache Relief by Time in Studies 4 and 5

	Study 4		Study 5		
	Placebo (n = 39)	Sumatriptan Injection 6 mg (n = 39)	Placebo (n = 88)	Sumatriptan Injection 6 mg (n = 92)	
Patients with pain relief (no/mild)					
5 Minutes post-injection	8%	21%	7%	23% ª	
10 Minutes post -injection	10%	49%³	25%	49%°	
15 Minutes post -injection	26%	74% ^a	35%	75% a	

P<0.05. n = Number of headaches treated An estimate of the cumulative probability of a patient with a cluster headache obtaining relief after being treated with either sumatriptan injection acebo is presented in Figure 1.



 $\label{thm:continuous} \begin{tabular}{ll} \textbf{The figure uses Kaplan-Meier (product limit) Survivorship Plot. Patients taking rescue medication were censored at 15 minutes. \end{tabular}$ The plot was constructed with data from patients who either experienced relief or did not require (request) rescue medication within a period of 2 hours following treatment. As a consequence, the data in the plot are derived from only a subset of the 258 headaches treated (rescue medication from only a subset of the 258 headaches treated (rescue medication from only a subset of the 258 headaches treated (rescue medication from only a subset of the 258 headaches treated (rescue medication from only a subset of the 258 headaches treated (rescue medication from only a subset of the 258 headaches treated (rescue medication from only a subset of the 258 headaches treated (rescue medication from only a subset of the 258 headaches treated (rescue medication from only a subset of the 258 headaches treated (rescue medication from only a subset of the 258 headaches treated (rescue medication from only a subset of the 258 headaches treated (rescue medication from only a subset of the 258 headaches treated (rescue medication from only a subset of the 258 headaches treated (rescue medication from only a subset of the 258 headaches treated (rescue medication from only a subset of the 258 headaches treated (rescue medication from only a subset of the 258 headaches treated (rescue medication from only a subset of the 258 headaches treated (rescue medication from only a subset of the 258 headaches treated (rescue medication from only a subset of the 258 headaches treated (rescue medication from only a subset of the 258 headaches treated (rescue medication from only a subset of the 258 headaches treated (rescue medication from only a subset of the 258 headaches treated (rescue medication from only a subset of the 258 headaches treated (rescue medication from only a subset of the 258 headaches treated (rescue medication from only a subset of the 258 headaches treated (rescue medication from only a subset of the 258 headaches treated (rescue medication from only a subset of the 258 headaches treated (rescue medication from only a $was \ required \ in \ 52 \ of \ the \ 127 \ placebo-treated \ headaches \ and \ 18 \ of \ the \ 131 \ headaches \ treated \ with \ sum a triptan \ injection).$ Other data suggest that treatment with sumatriptan injection is not associated with an increase in early recurrence of headache and has little effect on the incidence of later-occurring headaches (i.e., those occurring after 2, but before 18 or 24 hours).

Do not take more than 12 mg in a 24 hour period.

If you use too much sumatriptan injection, call you 16 HOW SUPPLIED/STORAGE AND HANDLING

Sumatriptan injection, USP contains sumatriptan (base) as the succinate salt and is supplied as a clear, colorless to pale yellow, sterile, nonpyrogenic solution as follows: NDC 43598-768-23 Sumatriptan Injection USP Autoinjector System includes 2 Autoinjectors, each with an associated single-dose

prefilled syringe which contains 6 mg of sumatriptan (as the succinate salt) and 3.5 mg of sodium chloride in 0.5 mL $\label{eq:store} \textbf{Store at 25°C (77°F); excursions permitted 15°C to 30°C (59°F to 86°F) [see USP Controlled Room Temperature]. Protect from light.}$ 17 PATIENT COUNSELING INFORMATION

 $\underline{Risk of Myocardial Ischemia and/or Infarction, Prinzmetal's Angina, Other Vasos pasm-Related Events, Arrhythmias, and Cerebrovascular Events}$ Inform patients that sumatriptan injection may cause serious cardiovascular side effects such as myocardial infarction or stroke. Although serious cardiovascular events can occur without warning symptoms, patients should be alert for the signs and symptoms of chest pain, shortness of breath, irregular heartbeat, significant rise in blood pressure, weakness, and slurring of speech, and should ask for medical advice if any indicative sign or $symptoms \ are \ observed. Apprise \ patients \ of the \ importance \ of this \ follow-up \ [see \textbf{Warnings} \ and \ Precautions \ (5.1, 5.2, 5.4, 5.5, 5.8)].$ Hypersensitivity Reactions

Inform patients that anaphylactic reactions have occurred in patients receiving sumatriptan injection. Such reactions can be life-threatening or fatal. In general, anaphylactic reactions to drugs are more likely to occur in individuals with a history of sensitivity to multiple allergens [see Contraindications (4), Warnings and Precautions (5.9)]. Concomitant Use with Other Triptans or Ergot Medications

Inform patients that use of sumatriptan injection within 24 hours of another triptan or an ergot-type medication (including dihydroergotamine or methysergide) is contraindicated [see Contraindications (4), Drug Interactions (7.1, 7.3)]. Serotonin Syndrome

Caution patients about the risk of serotonin syndrome with the use of sumatriptan injection or other triptans, particularly during combined use with SSRIs, SNRIs, TCAs, and MAO inhibitors [see Warnings and Precautions (5.7), Drug Interactions (7.4)]. Medication Overuse Headache

Inform patients that use of acute migraine drugs for 10 or more days per month may lead to an exacerbation of headache and encourage patients to record headache frequency and drug use (e.g., by keeping a headache diary) [see Warnings and Precautions (5.6)].

Advise patients to notify their healthcare provider if they become pregnant during treatment or plan to become pregnant [see Use in Specific Lactation

Ability to Perform Complex Tasks Treatment with sumatriptan injection may cause somnolence and dizziness; instruct patients administration of sumatriptan injection How to Use Sumatriptan Injection

they are able to self-administer sumatriptan injection in medically unsupervised situations. Instruct patients on storage and disposal of the pen [see Inform patients that the needle in the sumatriptan autoinjector penetrates approximately 1/4 of an inch (5 to 6 mm). Inform patients that the injection is intended to be given subcutaneously and intramuscular or intravascular delivery should be avoided. Instruct patients to use injection

PATIENT INFORMATION Sumatriptan (soo" ma trip' tan) Injection USP

What is the most important information I should know about sumatriptan injection Sumatriptan can cause serious side effects, including: Heart attack and other heart problems. Heart problems may lead to death

sites with an adequate skin and subcutaneous thickness to accommodate the length of the needle.

Stop taking sumatriptan and get emergency medical help right away if you have any of the following symptoms of a heart attack: discomfort in the center of your chest that lasts for more than a few minutes, or that goes away and comes back

- severe tightness, pain, pressure, or heaviness in your chest, throat, neck, or jaw pain or discomfort in your arms, back, neck, jaw, or stomach
- breaking out in a cold sweat nausea or vomiting feeling lightheaded Sumatriptan is not for people with risk factors for heart disease unless a heart exam is done and shows no problem. You have a higher risk for heart

shortness of breath with or without chest discomfort

- disease if you:

 have high blood pressure have high cholesterol levels
- are overweighthave diabetes have a family history of heart disea What is sumatriptan? Sumatriptan injection is a pres otion medicine used to treat acute migraine headaches with or without aura and acute cluster headaches in adults

Sumatriptan is not used to treat other types of headaches such as hemiplegic (that make you unable to move on one side of your body) or basilar (rare form of migraine with aura) migraines.

Sumatriptan is not used to prevent or decrease the number of migraine or cluster headaches you have It is not known if sumatriptan is safe and effective in children under 18 years of age.

- Do not take sumatriptan injection if you have: heart problems or a history of heart problems
- narrowing of blood vessels to your legs, arms, stomach, or kidneys (peripheral vascular disease)
- uncontrolled high blood pressure
- hemiplegic migraines or basilar migraines. If you are not sure if you have these types of migraines, ask your healthcare provide had a stroke, transient ischemic attacks (TIAs), or problems with your blood circulation
- taken any of the following medicines in the last 24 hours: o almotriptan (AXERT) o eletriptan (RELPAX) o frovatriptan (FROVA) o naratriptan (AMERGE) o rizatriptan (MAXALT, MAXALT-MLT)

o sumatriptan and naproxen (TREXIMET)

o ergotamines (CAFERGOT, ERGOMAR, MIGERGOT) o dihydroergotamine (D.H.E. 45, MIGRANAL) Ask your healthcare provider if you are not sure if your medicine is listed above.

- sumatriptan injection. $Before \ taking \ sum a triptan \ injection, tell \ your \ health care \ provider \ about \ all \ of \ your \ medical \ conditions, including \ if \ you \ medical \ conditions, including \ if \ you \ medical \ conditions, including \ if \ you \ medical \ conditions, including \ if \ you \ medical \ conditions, including \ if \ you \ medical \ conditions, including \ if \ you \ medical \ conditions, including \ if \ you \ medical \ conditions, including \ if \ you \ medical \ conditions, including \ if \ you \ medical \ conditions, including \ if \ you \ medical \ conditions, including \ if \ you \ medical \ conditions \ including \ if \ you \ medical \ conditions \ including \ if \ you \ medical \ conditions \ including \ if \ you \ medical \ conditions \ including \ if \ you \ medical \ conditions \ including \ if \ you \ medical \ conditions \ including \ if \ you \ medical \ conditions \ including \ if \ you \ medical \ conditions \ including \ if \ you \ medical \ conditions \ including \ if \ you \ medical \ conditions \ including \ if \ you \ medical \ conditions \ including \ if \ you \ medical \ conditions \ including \ if \ you \ medical \ conditions \ including \ if \ you \ medical \ including \ if \ you \ medical \ including \ including \ if \ you \ medical \ including \ including$
- have high cholesterol have diabetes
- have heart problems or family history of heart problems or stroke
- have kidney problems have liver problems
- have had epilepsy or seizures are not using effective birth control
- are breastfeeding or plan to breastfeed. Sumatriptan passes into your breast milk. It is not known if this can harm your baby. Talk with your healthcare provider about the best way to feed your baby if you take sumatriptan.
- Tell your healthcare provider about all the medicines you take, including prescription and over-the-counter medicines, vitamins, and herbal Sumatriptan and certain other medicines can affect each other, causing serious side effects.
- Especially tell your healthcare provider if you take antidepressant medicines called: selective serotonin reuptake inhibitors (SSRIs)
- serotonin norepinephrine reuptake inhibitors (SNRIs)
- tricyclic antidepressants (TCAs) monoamine oxidase inhibitors (MAOIs)
- $As kyour health care \, provider \, or \, pharmacist \, for \, a \, list \, of \, these \, medicines \, if you \, are \, not \, sure.$ $Know the \ medicines \ you \ take. \ Keep \ a \ list of them \ to \ show \ your \ healthcare \ provider \ or \ pharmacist \ when \ you \ get \ a \ new \ medicine.$
- How should I take sumatriptan injection? Certain people should take their first dose of sumatriptan injection in their healthcare provider's office or in another medical setting. Ask your
- healthcare provider if you should take your first dose in a medical setting. Use sumatriptan injection exactly as your healthcare provider tells you to use it.
- $\bullet \qquad \text{Your health care provider may change your dose}. Do not change your dose without first talking with your health care provider.}$ For adults, the usual dose is a single injection given just below the skin. You should give an injection as soon as the symptoms of your headache start, but it may be given at any time during a migraine or cluster
- headache attack. If you did not get any relief after the first injection, do not give a second injection without first talking with your healthcare provider. but not sooner.
- $\bullet \qquad \text{If you use too much sumatriptan injection, call your healthcare provider or go to the nearest hospital emergency room right away.}$ • You should write down when you have headaches and when you take sumatriptan injection so you can talk with your healthcare provider about
- how sumatriptan injection is working for you. What should I avoid while taking sumatriptan injection?

 $Sum a triptan \ can \ cause \ dizziness, \ weakness, \ or \ drows iness. \ If you have these \ symptoms, \ do not \ drive \ a \ car, \ use \ machinery, \ or \ do \ anything \ where \ you \ drows in \ drows \ drow$ need to be alert. What are the possible side effects of sumatriptan injection

 $Sum a triptan\ may\ cause\ serious\ side\ effects.\ See\ "What\ is\ the\ most\ important\ information\ I\ should\ know\ about\ sum a triptan\ injection?"$ These serious side effects include: changes in color or sensation in your fingers and toes (Raynaud's syndrome)

- stomach and intestinal problems (gastrointestinal and colonic ischemic events). Symptoms of gastrointestinal and colonic ischemic events o sudden or severe stomach pain o nausea or vomiting o stomach pain after meals o constipation or diarrhea o bloody diarrhea
- o weight loss problems with blood circulation to your legs and feet (peripheral vascular ischemia). Symptoms of peripheral vascular ischemia include:
- cramping and pain in your legs or hips feeling of heaviness or tightness in your leg muscles
- burning or aching pain in your feet or toes while resting numbness, tingling, or weakness in your legs cold feeling or color changes in 1 or both legs or feet
- medication overuse headaches. Some people who use too many sumatriptan injections may have worse headaches (medication overuse head a che). If your head a ches get worse, your health care provider may decide to stop your treatment with sum a triptant of the contract of the contract• serotonin syndrome. Serotonin syndrome is a rare but serious problem that can happen in people using sumatriptan injection, especially if
- sum a tript an injection is used with anti-depressant medicines called SSR Is or SNR Is.Call your healthcare provider right away if you have any of the following symptoms of serotonin syndrome mental changes such as seeing things that are not there (hallucinations), agitation, or coma
- changes in blood pressure
- high body temperature tight muscles
- trouble walking hives (itchy bumps); swelling of your tongue, mouth, or throat
- seizures. Seizures have happened in people taking sumatriptan injection who have never had seizures before. Talk with your healthcare provider about your chance of having seizures while you take sumatriptan injection The most common side effects of sumatriptan injection include:
- pain or redness at your injection site tingling or numbness in your fingers or toes
- dizziness warm, hot, burning feeling to your face (flushing)
- discomfort or stiffness in your neck feeling weak, drowsy, or tired
- Fell your healthcare provider if you have any side effect that bothers you or that does not go away. These are not all the possible side effects of sumatriptan injection. Call your doctor for medical advice about side effects. You may report side effects to FDA at 1-800-FDA-1088.
- How should I store sumatriptan injection? • Store at 25°C (77°F); excursions permitted 15°C to 30°C (59°F to 86°F)
- Store your medicine away from light. Keep your medicine in the packaging or carrying case provided with it. Keep sumatriptan injection and all medicines out of the reach of children.
- General information about the safe and effective use of sumatriptan injection Medicines are sometimes prescribed for purposes other than those listed in Patient Information leaflets. Do not use sumatriptan injection for a condition for which it was not prescribed. Do not give sumatriptan injection to other people, even if they have the same symptoms you have. It may
- This Patient Information leaflet summarizes the most important information about sumatriptan injection. If you would like more information, talk with your healthcare provider. You can ask your healthcare provider or pharmacist for information about sumatriptan injection that is written for healthcare professionals.
- For more information, call 1-888-375-3784 What are the ingredients in sum a tript an injection ?
- Inactive ingredients: sodium chloride, water for injection The other brands listed are trademarks of their respective owners and are not trademarks of Dr. Reddy's Laboratories Limited. The makers of these

INSTRUCTIONS FOR USE OF AUTOINJECTOR PEN

his device is called an Autoinjector pen. Here we use the shorter name 'pen'.

Important things that you need to know

brands are not affiliated with and do not endorse Dr. Reddy's Laboratories Limited or its products This Patient Information has been approved by the U.S. Food and Drug Administration.

SYSTEM Read this Patient Instructions for Use before you start to use Sumatriptan Autoinjector System. There may be new information. This information does not take the place of talking with your healthcare provider about your medical condition or treatment. You and your healthcare provider should talk about sumatriptan injection when you start taking it and at regular checkups.

SUMATRIPTAN INJECTION

INSTRUCTIONS FOR USE OF DISPOSABLE SUMATRIPTAN AUTOINJECTOR



Keep the Sumatriptan Autoinjector System out of the reach of children

1. Read all of the instructions carefully before using this pen. 2. Follow these step-by-step instructions every time you use the pen, as kyour doctor or pharmacist.3. Only use each pen once - do not try to use more than once. If you have any further questions.

A. ABOUT THE AUTO INJECTOR PEN The parts of the pen are shown in this picture.



B. GETTING READY Getting ready for the injection.

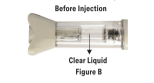
1. Wash your hands. 2. Choose an area with an adequate fatty tissue layer. 3. Clean the skin area to be injected with alcohol or a new sterile swab (see Figure A)



Getting the pen ready. 4. Take the pen out of the package.

5. Look in the transparent medicine window on the pen- Before injection, to check that the liquid is clear (see Figure B). If it is difficult to see what is in the window, hold the pen up to the light and check. • After injection, the blue plunger rod completely fills the medicine window (see Figure C).

• If the blue plunger rod can be seen through the medicine window, the device is spent and cannot be used again



6. Pull the gray cap off the pen.



7. Look inside the cap, check that the gray needle cover is inside (see Figure E).



8. Do not try to put the cap back . If you try to put it back, this will damage the needle ou are now ready to inject the medicine, go to step 9

C. INJECTING THE MEDICINE $. \ \ \textbf{Without pressing the blue button, push the pen firmly against your skin until you feel the stop point (see Figure F).}$ Pushing to the stop point unlocks the button. As long as the pen is firmly pressed against the skin, the safety lock is deactivated; the pen could fire unintentionally if the blue buttor is pressed by accident.

 Do not attempt to re-engage the safety lock at any time Keep the pen pressed against your skin for the next step:



 10. Keep pushing the pen against your skin then firmly press down the blue button on the top of the pen until it will not go further (see Figure G).
 You will hear a click, this indicates that the injection has started (see Figure G). • If the injection did not start, release the blue button, make sure the pen is pushed down against the skin and push down harder on the blue



Monitor the injection through the medicine window to make sure that the entire dose is injected. The blue plunger will move down the window, completely fill it, and stop moving when the injection is done (see Figure H).

• When the injection is done, keep holding the pen against the skin for 5 seconds (see Figure I). If you take the pen off before, not all of the





12. Carefully take the pen off your skin (see Figure J). The protective sleeve automatically covers the needle. It is then locked and the needle is protected



13. If you notice a spot of blood at the injection site, dab away with a cotton ball or tissue paper. Do not rub the injection site. If needed, you may cover the injection site with a bandage. $14. \ Visually \ check that \ there is \ no \ liquid \ left \ at \ the \ bottom \ of \ the \ medicine \ window. \ lifthere \ is \ liquid, consult \ your \ doctor \ or \ pharmacist.$

E. DISPOSE THE AUTOINJECTOR PEN 15. Discard the whole sumatriptan injection pen after use and discard the cap. Put your used injection pen in a FDA-cleared sharps disposal container right away after use (see Figure K).



. If you do not have a FDA-cleared sharps disposal container, you may use a household container that is

- o made of a heavy-duty plastic,
 o can be closed with a tight-fitting, puncture-resistant lid, without sharps being able to come out, upright and stable during use,
- o leak-resistant, and o properly labeled to warn of hazardous waste inside the containe

Issued: 12/2022

- When your sharps disposal container is almost full, you will need to follow your community guidelines for the right way to dispose of your sharps disposal container. There may be state or local laws about how you should throw away used needles and cartridges. For more information about safe sharps disposal, and for specific information about sharps disposal in the state that you live in, go to the FDA's website
- at: http://www.fda.gov/safesharpsdisposal. • Do not dispose of your used sharps disposal container in your h

 $Do \ not \ try \ to \ reuse \ the \ autoinjector \ pen. \ To \ avoid \ any \ injury, never \ try \ to \ touch \ the \ needle.$ This Patient Information and Instructions for Use has been approved by the U.S. Food and Drug Administration To reorder additional Patient information sheets contact Dr. Reddy's Customer Service at 1-866-733-3952.

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Dr.Reddy's

PHARMACIST - DETACH FROM HERE

tion site, dab away with ub the injection site. If swith a bandage. If at the bottom of the consult your doctor or