The International Journal of Neuropsychiatric Medicine

Unresolved Questions About Treatment-Resistant Anxiety Disorders

D.J. Stein and S. Seedat

Generalized Anxiety Disorder: Acute and Chronic Treatment

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Treatment-Resistant Posttraumatic Stress Disorder:
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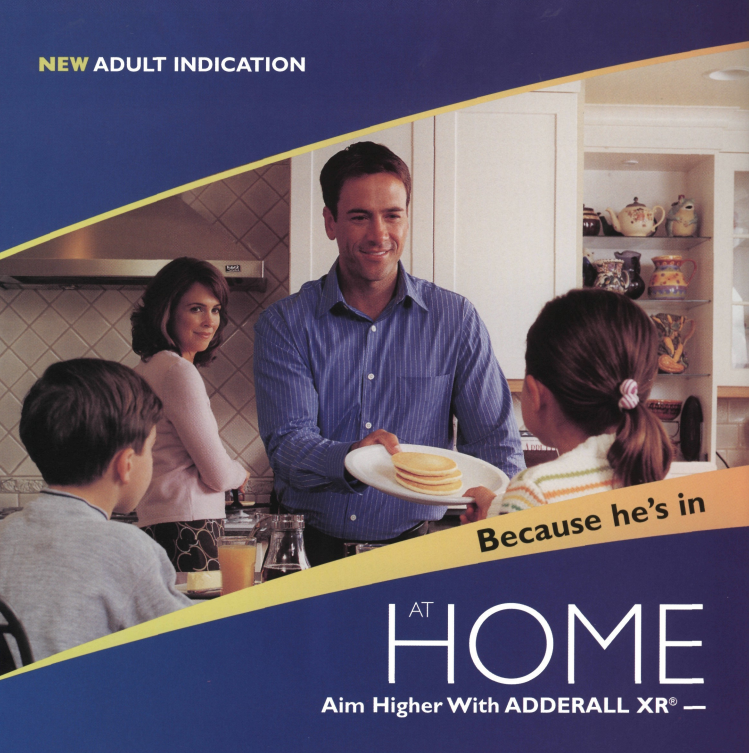
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Cognitive Deficits Following Coronary Artery Bypass Grafting: Prevalence, Prognosis, and Therapeutic Strategies

P.V. Raja, J.A. Blumenthal, and P.M. Doraiswamy

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The most common adverse events in pediatric trials included loss of appetite, insomnia, abdominal pain, and emotional lability. The most common adverse events in the adult trial included dry mouth, loss of appetite, insomnia, headache, and weight loss.

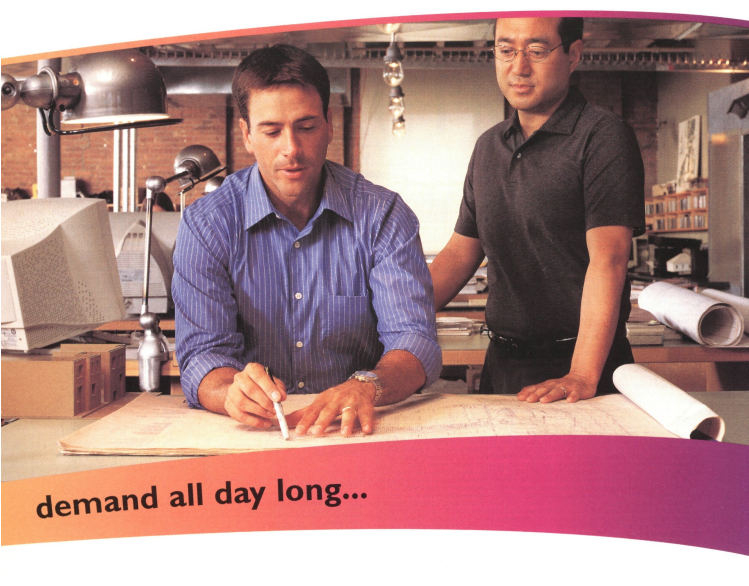
The effectiveness of ADDERALL XR for long-term use has not been systematically evaluated in controlled trials. As with other psychostimulants indicated for ADHD, there is a potential for exacerbating motor and phonic tics and Tourette's syndrome. A side effect seen with the amphetamine class is psychosis. Caution also should be exercised in patients with a history of psychosis.

Abuse of amphetamines may lead to dependence. Misuse of amphetamines may cause sudden death and serious cardiovascular adverse events. ADDERALL XR generally should not be used in children or adults with structural cardiac abnormalities. ADDERALL XR is contraindicated in patients with symptomatic cardiovascular disease, moderate to severe hypertension, hyperthyroidism and glaucoma, known hypersensitivity to this class of compounds, agitated states, history of drug abuse, or current or recent use of MAO inhibitors. ADDERALL XR should be prescribed with close physician supervision.

Reference: I. Data on file, Shire US Inc., 2002.

www.ADDERALLXR.com www.ADHDSupportCompany.com

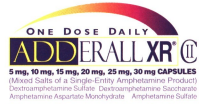
Please see brief summary of prescribing information on adjacent page.



WORK

For Efficacy That Measures Up to Life's Demands

- Once-daily dosing provides all-day symptom control
- · Mean ADHD-RS total scores for adults receiving ADDERALL XR decreased by 41%1
- ADDERALL XR is the only stimulant medication approved to treat adults with ADHD
- Clinical data in adults demonstrate that ADDERALL XR is generally well tolerated



Reach new heights

Shire

Shire US Inc.

...your ADHD support company™ 1-800-828-2088

ADDERALL XR® CAPSULES

CII Rx Only

ONE DOSE DAILY

5 mg, 10 mg, 15 mg, 20 mg, 25 mg, 30 mg CAPSULES

(Mixed Salts of a Single-Entity Amphetamine Product)
Dextroamphetamine Sulfate Dextroamphetamine Saccharate
Amphetamine Aspartate Monohydrate Amphetamine Sulfate

ERALL **XR**° @

AMPHETAMINES HAVE A HIGH POTENTIAL FOR ABUSE. ADMINISTRATION OF AMPHETAMINES FOR PROLONGED PERIODS OF TIME MAY LEAD TO DRUG DEPENDENCE. PARTICULAR ATTENTION SHOULD BE PAID TO THE POSSIBILITY OF SUBJECTS OBTAINING AMPHETAMINES FOR NON-THERAPEUTIC USE OR DISTRIBUTION TO OTHERS AND THE DRUGS SHOULD BE PRESCRIBED OR DISPENSED SPARINGLY.

MISUSE OF AMPHETAMINE MAY CAUSE SUDDEN DEATH AND SERIOUS CARDIOVASCULAR ADVERSE

INDICATIONS

ADDERALL XR® is indicated for the treatment of Attention Deficit Hyperactivity Disorder (ADHD). The efficacy of ADDERALL XR® in the treatment of ADHD was established on the basis of two controlled trials in children aged 6 to 12, and one controlled trial in adults who met DSM-1 criteria for ADHD (see CLINICAL PHARMACOLOGY), along with extrapolation from the known efficacy of ADDERALL9, the

formulation of this substance.

CONTRAINDICATIONS CONTAINDICATIONS
Advanced arteriosclerosis, symptomatic cardiovascular disease, moderate to severe hypertension, hyperthyroidism, known hypersensitivity or idiosyncrasy to the sympathominetic amines, glaucoma. Adjusted states. Patients with a history of drug abuse. During or within 14 days following the administration of monoamine oxidase inhibitors (hypertensive crises may result).

WARNINGS Wannings

Paychosis: Clinical experience suggests that, in psychotic patients, administration of amphetamine may exacerbate symptoms of behavior disturbance and thought disorder.

Long-Term Suppression of Growth: Data are inadequate to determine whether chron-

to use of stimulants in children, including amphetamine, may be causally associated with suppression of growth. Therefore, growth should be monitored during treatment, and patients who are not growing or gaining weight as expected should have their treatment interrupted.

Sudden Death and Pre-existing Structural Cardiac Abnormalities: Sudden death has been reported in association with amphetamine treatment at usual doses in children with structural cardiac abnormalities. Adderall XR® generally should not be used in children or adults with structural cardiac abnormalities.

PRECAUTIONS

General: The least amount of amphetamine feasible should be prescribed or dispensed at one time in order to

contract. The teast amount or amplementations reading to prescribed of dispensed at the limit in order to minimize the possibility of overdosage.

Hypertension: Caution is to be exercised in prescribing amphetamines for patients with even mild hypertension (see CONTRAINDICATIONS). Blood pressure and pulse should be monitored at appropriate intervals in patients taking ADDERALL XR®, especially patients with hypertension.

Tics: Amphetamines have been reported to exacerbate motor and phonic tics and Tourette's syndrome. Therefore, clinical evaluation for tics and Tourette's syndrome in children and their families should precede use of stimulant medications.

medications

Information for Patients: Amphetamines may impair the ability of the patient to engage in potentially hazardous activ ities such as operating machinery or vehicles; the patient should therefore be cautioned accordingly.

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Information for Patients: Amphetamines may impair the ability of the patient to engage in potentially hazardous activities such as operating machinery or vehicles; the patient should therefore be cautioned accordingly.

Drug Interactions: Aciditying agents—Gastrointestinal aciditying agents (guanethidine, reserptine, glutamic acid (Cl.), accorbic acid, etc.) (over absorption of amphetamines. University increasing university increases thereby increasing university (South microare) acid parts increase absorption of amphetamines. Ob-administration of ADDERALL XRP and gastrointestinal aikalinizing agents (southum bicarbonate, etc.), increase absorption of amphetamines. Ob-administration of ADDERALL XRP and gastrointestinal aikalinizing agents, such as antacids, should be avoided. Urinary aikalinizing agents (acetazoimalide, some thiazides) increase the concentration of the non-incirate species of the amphetamine molecule, thereby decreasing urinary excretion. Both groups of agents increase blood levels and therefore potentials the actions of amphetamines. Ardidepressants, ricycilo—Amphetamines may enhance the activity of through ambigressants or sympathonimines. Ardidepressants, increasing their effect on the release of nonephetamine metabolism. This slowing potentiates amphetamines university of the product of

Nonteratogenic Effects: Infants born to mothers dependent on amphetamines have an increased risk of premature delivery and low birth weight. Also, these infants may experience symptoms of withdrawal as demonstrated by dysphoria, including agitation, and significant lassitude.

Usage in Nursing Mothers: Amphetamines are excreted in human milk. Mothers taking amphetamines should be advised to refrain from nursing.

Pediatric Use: ADDERALL XR® is indicated for use in children 6 years of age and older.

Vesa in Children Under Six Years of Age: Effects of ADDERALL XR® in 3-5 year olds have not been studied. Long-term effects of amphetamines in children have not been well established. Amphetamines are not recommended for use in children under 3 years of age. Gerlatric Use: ADDERALL XR® has not been studied in the geriatric population.

ADVERSE EVENTS

ADVENSE EVEN IS

The premarketing development program for ADDERALL XR® included exposures in a total of 965 participants in clinical trials (635 pediatric patients, 248 adult patients, 82 healthy adult subjects). Of these, 635 patients (ages to 12) were evaluated in two controlled clinical studies, one open-label clinical study, and two single-dose clinical pharmacology studies (N=40). Safety data on all patients are included in the discussion that follows. Adverse

reactions were assessed by collecting adverse events, results of physical examinations, vital signs, weights, laboratory analyses, and ECGs.

Adverse events during exposure were obtained primarily by general inquiry and recorded by clinical investigators using terminology of their own choosing. Consequently, it is not possible to provide a meaningful estimate of the proportion of individuals experiencing adverse events without first grouping similar types of events into a smaller number of standardized event categories. In the tables and listings that follow, COSTART terminology has been used to classify reported adverse events

The stated frequencies of adverse events represent the proportion of individuals who experienced, at least once, a treatment-emergent adverse event of the type listed.

Adverse events associated with discontinuation of treatment: In two placebo-controlled studies of up to 5 weeks duration among children with ADHD, 2.4% (10/425) of ADDERALL XR® treated patients discontinued due to adverse events (including 3 patients with loss of appetite, one of whom also reported insomnia) compared to 2.7% (7/259) receiving placebo. The most frequent adverse events associated with discontinuation of ADDERALL XR® in controlled and uncontrolled, multiple-dose clinical trials of pediatric patients (N=595) are presented below. Over half of these patients were exposed to ADDERALL XR® for 12 months or more.

% of pediatric patients discontinuing (n=595)
2.9
1.5
1.2
1.0 Adverse event
Anorexia (loss of appetite)
Insomnia
Weight loss **Emotional lability** Depression 0.7

LIXR of the controlled 4-week study among adults with ADHD, patients who discontinued treatment due to adverse events among ADDERALL XR®-treated patients (N=191) were 3.1% (n=6) for nervousness including anxiety and irritability, 2.6% by Amphetamine Production and the controlled develope events among ADDERALL XR®-treated patients (N=191) were 3.1% (n=6) for nervousness including anxiety and irritability, 2.6% (n=5) for insomnia, 1% (n=2) each for headache, palpitation, and somnolence; and 0.5% (n=1) each for ALT increase, agitation, chest pain, cocalne craving, elevated blood pressure, and weight loss.

**Adverse events cocurring in a controlled trial: Adverse events reported in a 3-week clinical trial in adults treated with ADDERALL XR® or placebo are presented in the tables below.

presented in the labels below.

The prescriber should be aware that these figures cannot be used to predict the incidence of adverse events in the course of usual medical practice where patient characteristics and other factors differ from those which prevailed in the clinical trials. Similarly, the cited frequencies cannot be compared with figures obtained from other clinical investigations involving different treatments, uses, and investigations. The cited figures, however, do provide the prescribing physician with some basis for estimating the relative contribution of drug and non-drug factors to the adverse event incidence rate in the population studied.

Table 1 Adverse Events Reported by More Than 1% of Pediatric Patients Receiving ADDERALL XR® with

Body System	Preferred Term	ADDERALL XR® (n=374)	Placebo (n=210)
General	Abdominal Pain (stomachache)	14%	10%
	Accidental Injury	3%	2%
	Asthenia (fatique)	2%	0%
	Fever	5%	2%
	Infection	4%	2%
	Viral Infection	2%	0%
Digestive	Loss of Appetite	22%	2%
System	Diarrhea	2%	1%
	Dyspepsia	2%	1%
	Nausea	5%	3%
	Vomiting	7%	4%
Nervous System	Dizziness	2%	0%
	Emotional Liability	9%	2%
	Insomnia	17%	2%
	Nervousness	6%	2%
Metabolic/Nutritional	Weight Loss	4%	0%

Table 2 Adverse Events Reported by 5% or More of Adults Receiving ADDERALL XR® with Higher Incidence Than on Placebo in a 255 Patient Clinical Forced Weekly-Dose Titration Study*

Body System	Preferred Term	ADDERALL XR® (n=191)	Placebo (n=64)
General	Asthenia Headache	6% 26%	5% 13%
Digestive System	Loss of Appetite Diarrhea Dry Mouth Nausea	33% 6% 35% 8%	3% 0% 5% 3%
Nervous System	Agitation Anxiety Dizziness Insomnia	8% 8% 7% 27%	5% 5% 0% 13%
Cardiovascular System	Tachycardia	6%	3%
Metabolic/Nutritional	Weight Loss	11%	0%
Urogenital System	Urinary Tract Intection	5%	0%

Note: The following events did not meet the criterion for inclusion in Table 2 but were reported by 2% to 4% of adult patients receiving ADDERALL XRP with a higher incidence than patients receiving placebo in this study infection, photosensitivity reaction, constipation, toolth disorder, emotional lability, libido decreased, somnolence, speech disorder, palpitation, twitching, dyspnea, sweating, dysmenorrhea, and impotence.

included doses up to 60 mg.

The following adverse reactions have been associated with amphetamine use: Cardiovascular: Palpitations, tachycardia, elevation of blood pressure, sudden death, myocardial infarction. There have been isolated reports cardiomyogathy associated with chronic amphetamine use. Central Nervous System: Psychotic episodes at recommended doses, overstimulation, restlessness, dizziness, insomnia, euphoria, dyskinesia, dysphoria, depression, tremor, headache, exacerbation of motor and phonic tics and Tourette's syndrome, seizures, stroke. Gastrointestinal: Dryness of the mouth, unpleasant taste, diarrhea, constipation, other gastrointestinal disturbances. Anorexia and weight loss may occur as undesirable effects. Allergic: Urticaria. Endocrine: Impotence, changes in libido.

DRUG ABUSE AND DEPENDENCE

ADDERALL XR® is a Schedule II controlled substance

ADDEFABLL XA** is a Schedule if controlled substance. Amphetamines have been extensively abused. Tolerance, extreme psychological dependence, and severe social disability have occurred. There are reports of patients who have increased the dosage to many times that recommended. Abrupt cessation following prolonged high dosage administration results in extreme fatigue and mental depression; changes are also noted on the sleep EEG. Manifestations of chronic intoxication with amphetamines may include severe dermatoses, marked insomnia, irritability, hyperactivity, and personality changes. The most severe manifestation of chronic intoxication is psychosis, often clinically indistinguishable from schizophrenia.

OVERDOSAGE

OVERDOSAGE
Individual patient response to amphetamines varies widely. Toxic symptoms may occur idiosyncratically at low doses. Symptoms: Manifestations of acute overdosage with amphetamines include restlessness, tremor, hyper-reflexia, rapid respiration, confusion, assaultiveness, hallucinations, panic states, hyperoyrexia and rhabdomyosyis. Fatigue and depression usually follow the central nervous system stimulation. Cardiovascular effects include arrhythmias, hypertension or hypotension and circulatory collapse. Gastrointestinal symptoms include nausea, vomiting, diarrhea, and abdominal cramps. Fatal poisoning is usually preceded by convulsions and coma. Treatment: Consult with a Certified Poison Control Center for up to date guidance and advice. Management of acute amphetamine intoxication is largely symptomatic and includes gastric lavage, administration of activated charcoal, administration of a cathartic and sedation. Experience with hemodialysis or peritoneal dialysis is inadequate to permit recommendation in this regard. Acidification of the urine increases amphetamine excretion, but is believed to increase risk of acute renal failure if myoglobinuria is present. If acute severe hypertension complicates amphetamine overdosage, administration of intravenous phentolamine has been suggested. However, a gradual drop in blood pressure will usually result when sufficient sedation has been achieved. Chlorpromazine antagonizes the central stimulant effects of amphetamines and can be used to treat amphetamine intoxication.

The prolonged release of mixed amphetamines as from ADDERALL XR® should be considered when treating

The prolonged release of mixed amphetamine salts from ADDERALL XR® should be considered when treating patients with overdose.

Dispense in a tight, light-resistant container as defined in the USP. Store at 25° C (77° F). Excursions permitted to 15-30° C (59-86° F) [see USP Controlled Room Temperature].

Manufactured for: Shire US Inc., Newport, KY 41071 Made in USA For more information call 1-800-828-2088, or visit www.adderallxr.com. ADDERALL® and ADDERALL XR® are registered in the US Patent and Trademark Office. Copyright ©2004 Shire US Inc.

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Lisa Arrington Shoshana Bauminger The Anxiety Disorders
Association of America
announces its

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Available online at www.adaa.org

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JUNIOR FACULTY RESEARCH GRANT - \$30,000 Deadline: Tuesday, December 9, 2004

TRAINEE TRAVEL AWARD - \$1,500 Deadline: Monday, December 20, 2004

Fifteen awardees are selected to attend the ADAA's 25th Annual Conference, March 17-20, 2005, in Seattle, Washington. To date, the ADAA Awards Program has given more than 75 travel awards and 15 research grants, totaling nearly \$700,000.

For award descriptions, eligibility requirements, award criteria, and applications, please visit the ADAA Web site at www.adaa.org. For more information, contact the ADAA Awards program manager, Jane Caroline Parham, at (P) 240-485-1016, (F) 240-485-1035, or email at jparham@adaa.org.

About the ADAA

The ADAA is the only national, nonprofit partnership of researchers, health care professionals, and individuals dedicated solely to the early diagnosis, prevention, and treatment of anxiety disorders. It is the Association's goal to promote professional and public awareness and understanding of anxiety disorders. It also seeks to increase the availability of effective treatment, reduce the stigma surrounding anxiety disorders, and stimulate research.



Anxiety Disorders Association of America 8730 Georgia Avenue, Suite 600 Silver Spring, MD 20910, USA Phone: 240-485-1001 Web site: www.adaa.org BRIEF SUMMARY of PRESCRIBING INFORMATION

RIDICATIONS AND USAGE: Bipdar Mania: SERDOUEL is indicated for the short-term treatment of acute manic
episodes associated with bipdar if disorder, as either monotherapy or adjunct therapy to lithium or divelprose. The
efficacy of SERDOUEL in acute bipdar mania was established in two 3-week monotherapy trials and one 3-week
dipunct therapy trial of bipdar is platient initially hospitated for up to 7 days for acute manic Effectiveness for
more than 3 weeks has not been systematically evaluated in clinical trials. Therefore, the physician who elects to use
SERDOUEL for extended periods should periodically re-evaluate the long-term issis and benefits of the drug for
the individual patient. Serboughemants: SERDOUEL is indicated for the treatment of schizophrenia, the efficacy of
REPOLICE in orderpolare was established in short-them (fewerk) controlled trials of schizophrenia was established in short-them (fewerk) controlled trials. Therefore, the physician who elects to use SERDOUEL for extended periods should
periodically re-evaluate the long-term usefulness of the drug for the individual patient.

periodically re-evaluate the long-term usefulness of the drug for the individual patient.

CONTRAINDICATIONS: SEROOUEL is contraindicated in individuals with a known hypersensitivity to this medication or any of its ingredients.

WARHINGS: Neurologic Realizemal Syndrems (NMS): A potentially fatal symptom complex sometimes referred to as Neurologic Malignant Syndroms (NMS) has been reported in association with administration of artipsychotic drugs, including SEROOUEL. Bare cases of NMS have been reported with SEROOUEL clinical manifestations of the NMS are hyperyreain, unsaler inglish. Better interest states, and evidence of automatic instability irregular pulse or blood pressure, bathyoraria, displorerses, and carriace dysriptiminal. Additional signs may include elevated creative prospectives, including states of the Malignant Syndrome is complicated. In artificial and adaptives, it is important to exclude cases where the clinical presentation includes both seroise medical lines (e.g., presentional, systemic infection, etc.) and untrasted or inadequately retailed excrapyramidal signs and symptoms (EPS). Other important considerations in the differential diagnosis include carrial anticipation of the code, heat stroke, and plant ever and primary contribution of antipsychotic drugs and other contributions of antipsychotic drugs and other contributions of any concomitant serious medical problems for which specific teatments are available. There is no general agreement about specific pharmacological bendament enginess for MMS, in a plant approach of a carrially considered. In a plant and advantage of the provision of the posterior and approach of the posterior and problems are advantaged. The proposition of the posterior and problems are advantaged to a carrially considered. The production of the posterior excertions of the posterior and problems are advantaged to a carrially considered. The production of the posterior excertions of the posterior and problems are advantaged to a carrially reconsidered. The residence

periodically during treatment. Any patient treated with altypical artispsychotics should be monitored for symptoms of hyper-pylorenia during treatment with altypical artispsychotics should undergo basing blood glucose testing. In some cases, proportyperam is not received with the applical artispsychotic was discontinuation of anti-clarbent treatment despite discontinuation of the suspect drug.

PRECAUTIONS: Caemaric Ordinactive Hypothesianics: ESPOLUEL may induce orthostatic hypotension associated with disciness, bachyractic and, in some patients, syncope, especially during the initial dose-initiation period, probably reflecting its, or advancer patients and in the patients treated with SEROOUEL, compared with D% (0007) on placebo and about 0.4% (2627) on active control drugs. SEROOUEL, sold be used with particular catation in plants with howar cardiovascular disease, lichary or importantial infarction or submine heart diseases, heart failure or conduction abnormabiles, perservoscular diseases. Patent failure or conduction abnormabiles, perservoscular diseases or conditions with or would preclapose patients on the potential patients with north conditions. The risk of orthrestatic hypotension and synopose may be minimized by limiting the intuition schoolule in appropriate character. The development of extractive was observed in seasociation with a struction schoolule in appropriate character. The development of extractive was observed in seasociation with a struction schoolule in a propriate character. The development of extractive was observed in seasociation with a structive character. The development of extractive was observed in seasociation with a structive character. The development of extractive was observed in patients with a structive character. The propriate in any positive patients with a structive character of the patients. The propriate in a popular patients was an about the analysis of the patients. The propriate in any positive patients with a structure of the patients when the patients and patien

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Steven Johnson syndrome (SJS).

DRUB ABLISE AND DEPENDENCE: Controlled Substance Class: SERCOULE, is not a controlled substance. Physicial and Psychological dependence: SERCOULE, has not been systematically studied, in animals or humans, for its potential for abuse, tolerance or physicial dependence. While the climical trials did not reveal any tendency for any drug-selding behavior, these observations were not systematic and it is not possible to predict on the basic of this limit et experience the start to which a OSF-active drug will be missed, diverted, and/or abused once to extend to which a OSF-active drug will be missed, diverted, and/or abused once to extend carefully for a history of drug abuse, and such patients should be observed dozely for syster of misses or abuse of SERCOULE, e.g., development of tolerance, increases in dose, drug-seeling behavior.

observed dosely for signs of misuse or abuse of SERVOUEL, e.g., overlopment of tolerance, increases in dose, unjus-seking betavor.

OVERDOSAGE: Numae appriames: Experience with SEROQUEL (quistignine furmarte) in acute overdosage and in fatalities. In general, reported signs and symptoms were those resulting from an exaggeration of the drugs known pharmacological effects is, e. roscinences and soldant, backyvartia and hypotension. One case, involving an estimated overdose of SEROQUEL some resulting indept book in post-marting operations, there have been very rare reports of overdoses of SEROQUEL alone resulting in death, come or OTc prolongation. Management of Overdosage, in case of acute overdosage, establish and maintain an airway and ensure adequate oxygenation and verificition. Sective of those oxygenation and verificition. Sective of videos oxygenation and verificition. Sective of videos oxygenation and verificition. Sective of videos overdosage, setablish and maintain an airway and ensure adequate oxygenation and verificition. Sective of videos oxygenation and verificition. Sective of videos oxygenation and verificition. Sective of videos overdosage oxygenation and verificition. Sective of videos overdosage oxygenation and verificition. Sective of videos overdosage oxygenation and verificition. Sective of videos videosage oxygenation and verificition. Sective of videos videosage oxygenation and verificition. Sective oxygenation videosage oxygenation and very oxygenatic sections oxygenation and very oxygenatic videosage oxygenatic videosage oxygenation and very oxygenatic videosage oxygenation and very oxygenatic videosage oxygenation and very oxygenatic videosage oxygenation very oxygenatic videosage oxygenation very

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SEROQUEL is indicated for the treatment of acute manic episodes associated with bipolar I disorder and the treatment of schizophrenia. Patients should be periodically reassessed to determine the need for continued treatment.

Prescribing should be consistent with the need to minimize the risk of tardive dyskinesia, seizures, and orthostatic hypotension. A rare condition referred to as neuroleptic malignant syndrome (NMS) has been reported with this class of medications, including SEROQUEL.

There have been reports of diabetes mellitus and hyperglycemia-related adverse events associated with the use of atypical antipsychotics, including SEROQUEL.

The most commonly observed adverse events associated with the use of SEROQUEL in clinical trials were somnolence, dry mouth, dizziness, constipation, asthenia, abdominal pain, postural hypotension, pharyngitis, SGPT increase, dyspepsia, and weight gain.





AstraZeneca Pharmaceuticals LP

To prevent medication errors, write "SEROQUEL" clearly on your Rx pad. Spell "SEROQUEL" clearly over the phone. **Redefine Success**

www.SEROQUEL.com Please see Brief Summary of Prescribing Information on following page.

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The International Journal of Neuropsychiatric Medicine

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EDITORIAL MISSION

CNS Spectrums' editorial mission is to address relevant neuropsychiatric topics, including the prevalence of comorbid diseases among patients, and original research and reports that emphasize the profound diagnostic and physiologic connections made within the neurologic and psychiatric fields. The journal's goal is to serve as a resource to psychiatrists and neurologists seeking to understand and treat disturbances of cognition, emotion, and behavior as a direct consequence of central nervous system disease, illness, or trauma.

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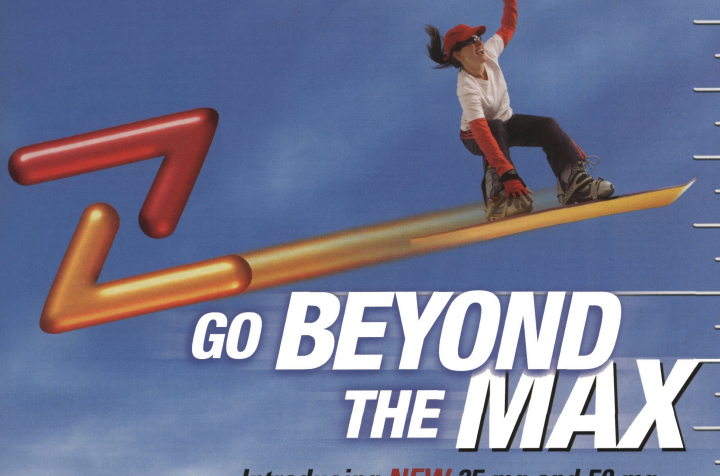
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Introducing **NEW 25** mg and 50 mg capsules of **ZONEGRAN®** (zonisamide)



ZONEGRAN is indicated as adjunctive therapy in the treatment of partial seizures in adults with epilepsy.

In clinical trials, the most common adverse events that occurred with ZONEGRAN were somnolence, dizziness, anorexia, headache, nausea, and agitation/irritability.

*Can also be dosed twice daily.

Please see brief summary of Prescribing Information on adjacent page.

References: 1. ZONEGRAN® Prescribing Information. Elan Pharmaceuticals. 2002. 2. Brodie M, Wilson E, Smith D, et al. Steady-state drug interaction study of zonisamide and lamotrigine in epileptic patients. Neurology. 2001;56(3):A337 (abstract). 3. Data on file. Elan Pharmaceuticals, Inc.



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More dosing options for meeting patients' needs

- Increase your dosing flexibility
- Choose from 3 dosage strengths:
 25 mg, 50 mg, and 100 mg capsules
- Tailor therapy to the individual patient

Proven efficacy with confidence-building benefits¹⁻³

- Few drug-to-drug interactions
- Minimal cognitive impairment
- 63-hour half-life—the longest of any newer AED
- Convenient QD dosing*



CONTRAINDICATIONS

ZONEGRAN is contraindicated in patients who have demonstrated hypersensitivity to sulfonamides or zonisamide.

WARNINGS

Potentially Fatal Reactions to Sulfonamides: Fatalities have occurred, although rarely, as a result of severe reactions to sulfonamidel (Zonisamide is a sulfonamide) including Stevens-Johnson syndrome, toxic epidermal necrolysis, fulminant hepatic necrosis, agranulocytosis, aplastic anemia, and other blood dyscrasias. Such reactions may occur when a sulfonamide is readministered irrespective of the route of administration. It signs of hypersensitivity or other serious reactions occur, discontinue zonisamide immediately. Specific experience with sulfonamide-type adverse reaction to zonisamide is described below.

below.

Serious Skin Reactions: Consideration should be given to discontinuing ZONEGRAN in patients who develop an otherwise unexplained rash. If the drug is not discontinued, patients should be observed frequently. Seven deaths from severe rash [i.e. Stevens-Johnson syndrome (SJS) and toxic epidermal necrolysis (TENI) were reported in the first 11 years of marketing in Japan. All of the patients were receiving other drugs in addition to zonisamide. In post-marketing experience from Japan, a total of 49 cases of SJS or TEN have been reported, a reporting rate of 46 per million patient-years of exposure. Although this rate is greater than background, it is probably an underestimate of the true incidence because of under-reporting. There were no confirmed cases of SJS or TEN in the US, European, or Japanese development programs.

In the US and European randomized controlled trials, 6 of 269 (2.2%) zonisamide patients/discontinued treatment because of rosh compared to none on placebo. Across all trials during the US and European development, rash that led to discontinuation of zonisamide was reported in 1.4% of patients (12.0 events per 1000 patient-years of exposure). During Japanese development, serious rash or rash that led to study drug discontinuation was reported in 2.0% of patients (27.8 events per 1000 patient years). Rash usually occurred early in treatment, with 85% reported within 16 weeks in the US and European studies and 90% reported within two weeks in the Japanese studies. There was no apparent relationship of dose to the occurrence of rash.

Serious Hematologic Events: Two confirmed cases of aplastic anemia and one confirmed case of agranulocytosis were reported in the first 11 years of marketing in Japan, rates greater than generally accepted background rates. There were no cases of aplastic anemia and two confirmed cases of agranulocytosis in the US, European, or Japanese development programs. There is inadequate information to assess the relationship, if any, between dose and duration of treatment and these events.

Oligohidrosis and Hyperthermia in Pediatric Patients:

Oligohidrosis, sometimes resulting in heat stroke and hospitalization, is seen in association with zonisamide in pediatric patients.

During the pre-approval development program in Japan, one case of oligohidrosis was reported in 403 pediatric patients, an incidence of 1 case per 285 patient-years of exposure. While there were no cases reported in the US or European development programs, fewer than 100 pediatric patients participated in these trials.

In the first 11 years of marketing in Japan, 38 cases were reported, an estimated reporting rate of about 1 case per 10,000 patient-years of exposure. In the first year of marketing in the US, 2 cases were reported, an estimated reporting rate of about 12 cases per 10,000 patient-years of exposure. These rates are underestimates of the true incidence because of under-reporting. There has also been one report of heat stroke in an 18-year-old patient in the US.

Decreased sweating and an elevation in body temperature above normal characterized these cases. Many cases were reported after exposure to elevated environmental temperatures. Heat stroke, requiring hospitalization, was diagnosed in some cases. There have been no reported deaths.

cases. There have been no reported aceams.

Pediatric patients appear to be at an increased risk for zonisamide-associated oligohidrosis and hyperthermica. Patients, especially pediatric patients, treated with Zonegran should be monitored closely for evidence of decreased sweating and increased body temperature, especially in warm or hot weather. Caution should be used when zonisamide is prescribed with other drugs that predispose potients to hear-related disorders; these drugs include, but are not limited to, carbonic anhydrase inhibitors and drugs with anticholinergic activity.

The practitioner should be aware that the safety and effectiveness of zonisamide in pediatric patients have not been established, and that zonisamide is not approved for use in pediatric patients.

Seizures on Withdrawal: As with other AEDs, abrupt withdrawal of ZONEGRAN in patients with epilepsy may precipitate increased seizure frequency or status epilepticus. Dose reduction or discontinuation of zonisamide should be done gradually.

or discontinuation of zonísamide should be done gradually.

Teratogenicity: Women of child bearing potential who are
given zonisamide should be advised to use effective contraception. Zonisamide was teratogenic in mice, rats, and dags and
embryolethal in monkeys when administered during the period
of organogenesis. A variety of fetal abnormalities, including
cardiovascular defects, and embryo-fetal deaths occurred at
maternal plasma levels similar to or lower than therapeutic
levels in humans. These findings suggest that the use of ZONEGRAN during pregnancy in humans may present a significant
risk to the fetus (see PRECAUTIONS, Pregnancy subsection). It
cannot be said with any confidence, however, that even mild
seizures do not pose some hazards to the developing fetus.
Zonisamide should be used during pregnancy only if the potential benefit justifies the potential risk to the fetus.

Coantitye/ Neuropsychiatric Adverse Events: Use of ZONE-

Cognitive/ Neuropsychiatric Adverse Events: Use of ZONE-GRAN was frequently associated with central nervous systemrelated adverse events. The most significant of these can be classified into three general categories: 1) psychiatric symptoms, including depression and psychosis, 2) psychomotor slowing, difficulty with concentration, and speech or language problems, in particular, word-finding difficulties, and 3) somnolence or fatigue.

nolence or tatigue.

In placebo-controlled trials, 2.2% of patients discontinued ZONEGRAN or were hospitalized for depression compared to 0.4% of placebo patients, while 1.1% of ZONEGRAN and 0.4% of placebo patients attempted suicide. Among all epilepsy patients treated with ZONEGRAN, 1.4% were discontinued and 1.0% were hospitalized because of reported depression suicide attempts. In placebo-controlled trials, 2.2% of patients discontinued ZONEGRAN or were hospitalized due to psychosis or psychosis-related symptoms compared to none of the placebo potients. Among all epilepsy patients treated with ZONEGRAN, 0.9% were discontinued and 1.4% were hospitalized because of reported psychosis or related symptoms.

Psychomotor slowing and difficulty with concentration occurred in the first month of treatment and were associated with doses above 300 mg/day. Speech and language problems tended to occur after 6–10 weeks of treatment and at doses above 300 mg/day. Although in most cases these events were of mild to moderate severity, they at times led to withdrawal from treatment.

Somnolence and fatigue were frequently reported CNS adverse events during clinical trials with ZONEGRAN. Although in most cases these events were of mild to moderate severity, they led to withdrawal from treatment in 0.2% of the patients enrolled in controlled trials. Somnolence and fatigue tended to occur within the first month of treatment. Somnolence and to tigue occurred most frequently at doses of 300–500 mg/day. Patients should be cautioned about this possibility and special care should be taken by patients if they drive, operate machinery, or perform any hazardous task.

PRECAUTIONS

General: Somnolence is commonly reported, especially at higher doses of ZONEGRAN (see WARNINGS: Cognitive/Neuropsychiatric Adverse Events subsection). Zonisamide is metabolized by the liver and eliminated by the kidneys caution should therefore be exercised when administering ZONEGRAN to patients with hepatic and renal dysfunction (see CLINICAL PHARMACOLOGY, Special Populations subsection of full Precribing Information).

see CLINICAL PRAKMACOLOGY, Special Populations subsection of full Precribing Information).

Kidney Stones: Among 991 patients treated during the development of ZONEGRAN, 40 patients (4.0%) with epilepsy receiving ZONEGRAN developed clinically possible or confirmed kidney stones (e.g. clinical symptomatology, sonography, etc.), a rate of 34 per 1000 patienty-vears of exposure). Of these, 12 were symptomatic, and 28 were described as possible kidney stones based on sonographic detection. In nine patients, the diagnosis was confirmed by a passage of a stone or by a definitive sonographic finding. The rate of occurrence of kidney stones was 28.7 per 1000 patienty-ears of exposure in the first six months, 62.6 per 1000 patienty-ears of exposure in the first six months, 62.6 per 1000 patienty-ears of exposure ofter 12 months of use. There are no normative sonographic data available for either the general population or patients with epilepsy. The clinical significance of the sonographic finding is unknown. The analyzed stones were composed of calcium or urate salts. In general, increasing fluid intake and urine output can help reduce the risk of stone formation, particularly in those with predisposing risk factors. It is unknown, however, whether these measures will reduce the risk of stone formation in patients treated with ZONEGRAN.

Effect on Renal Function: In several clinical studies, zonisamide

in patients treated with ZONEGRAN.

Effect on Renal Function: In several clinical studies, zonisamide was associated with a statistically significant 8% mean increase from baseline of serum creatinine and blood urea nitrogen (BUN) compared to essentially no change in the placebo patients. The increase appeared to persist over time but was not progressive; this has been interpreted as an effect on glomerular filtration rate (GFR). There were no episodes of unexplained acute renal failure in clinical development in the US, Europe, or Japan. The decrease in GFR appeared within the first 4 weeks of treatment. In a 30-day study, the GFR returned to baseline within 2–3 weeks of drug discontinuation. There is no information about reversibility, after drug discontinuation, of the effects on GFR after long-term use. ZONEGRAN should be discontinued in patients who develop acute renal failure or a clinically significant sustained increase in the creatinine/BUN concentration. ZONEGRAN should not be used in patients with renal failure (estimated GFR < 50 mL/min) as there has been insufficient experience concerning drug dosing and toxicity.

insufficient experience concerning drug dosing and toxicity.

Sudden Unexplained Death in Epilepsy: During the development of ZONEGRAN, nine sudden unexplained deaths occurred among 991 patients with epilepsy receiving ZONE-GRAN for whom accurate exposure data are available. This represents an incidence of 7.7 deaths per 1000 patient years. Although this rate exceeds that expected in a healthy population, it is within the range of estimates for the incidence of sudden unexplained deaths in patients with refractory epilepsy not receiving ZONEGRAN (ranging from 0.5 per 1000 patient-years for the general population of patients with reliplepsy, to 2–5 per 1000 patient-years for patients with refractory epilepsy; higher incidences range from 9–15 per 1000 patient-years among surgical candidates and surgical failures). Some of the deaths could represent seizure-related deaths in which the seizure was not observed.

Status Epilepticus: Estimates of the incidence of treatment emergent status epilepticus in ZONEGRAN-treated patients are difficult because a standard definition was not employed. Nonetheless, in controlled trials, 1.1% of patients treated with ZONEGRAN had an event labeled as status epilepticus compared to none of the patients treated with placebo. Among patients treated with ZONEGRAN across all epilepsy studies (controlled and uncontrolled), 1.0% of patients had an event reported as status epilepticus.

Creatine Phosphokinase (CPK) Elevation and Pancreatitis: In the post-market setting, the following rare adverse events have been observed (<1:1000):

If patients taking zonisamide develop severe muscle pain

and/or weakness, either in the presence or absence of a fever, markers of muscle damage should be assessed, including serum CPK and aldolase levels. If elevated, in the absence of another obvious cause such as traumo, grand mal seizures, etc., topering and/or discontinuance of zonisamide should be considered and appropriate treatment initiated.

Patients taking zonisamide that manifest clinical signs and symptoms of pancreatitis should have pancreatic lipase and amylase levels monitored. If pancreatitis is evident, in the absence of another obvious cause, tapering and/or discontinuation of zonisamide should be considered and appropriate treatment initiated.

Information for Patients: Patients should be advised as follows:

- 7. ZONEGRAN may produce drowsiness, especially at higher doses. Patients should be advised not to drive a car or operate other complex machinery until they have gained experience on ZONEGRAN sufficient to determine whether it affects their performance.
- Patients should contact their physician immediately if a skin rash develops or seizures worsen.
- 3. Patients should contact their physician immediately if they develop signs or symptoms, such as sudden back pain, abdominal pain, and/or blood in the urine, that could indicate a kidney stone. Increasing fluid intake and urine output may reduce the risk of stone formation, particularly in those with predisposing risk factors for stones.
- Patients should contact their physician immediately if a child has been taking ZONEGRAN and is not sweating as usual with or without a fever.
- Because zonisamide can cause hematological complications, patients should contact their physician immediately if they develop a fever, sore throat, oral ulcers, or easy bruising.
- 6. As with other AEDs, patients should contact their physician if they intend to become pregnant or are pregnant during ZONEGRAN therapy. Patients should notify their physician if they intend to breast-feed or are breast-feeding an infant.
- Patients should contact their physician immediately if they develop severe muscle pain and/or weakness.

Laboratory Tests: In several clinical studies, zonisamide was associated with a mean increase in the concentration of serum creatinine and blood urea nitrogen (BUN) of approximately 8% over the baseline measurement. Consideration should be given to monitoring renal function periodically (see PRECAUTIONS, Effect on Renal Function subsection).

Zonisamide was associated with an increase in serum alkaline phosphatase. In the randomized, controlled trials, a mean increase of approximately 7% over boseline was associated with zonisamide compared to a 3% mean increase in placebo-treated potients. These changes were not statistically significant. The clinical relevance of these changes is unknown.

Drug Interactions: Effects of ZONEGRAN on the pharmacokinetics of other antiepilepsy drugs (AEDs): Zonisamide had no appreciable effect on the steady state plasma concentrations of phenytoin, carbamazepine, or valproate during clinical trials. Zonisamide did not inhibit mixed-function liver oxidase nezymes (cytochrome P450), as measured in human liver microsomal preparations, in vitro. Zonisamide is not expected to interfere with the metabolism of other drugs that are metabolized by cytochrome P450 isozymes.

Effects of other drugs on ZONEGRAN pharmacokinetics: Drugs that induce liver enzymes increase the metabolism and clearance of zonisamide and decrease its half-life. The half-life of zonisamide following a 400 mg dose in patients concurrently on enzyme-inducing AEDs such as phenytoin, carbamazepine, or phenobarbital was between 27–38 hours; the half-life of zonisamide in patients concurrently on the non-enzyme inducing AED, valproate, was 46 hours. Concurrent medication with drugs that either induce or inhibit CYP3A4 would be expected to alter serum concentrations of zonisamide.

Interaction with cimetidine: Zonisamide single dose pharmacokinetic parameters were not affected by cimetidine (300 mg four times a day for 12 days).

Carcinogenicity, Mutagenesis, Impairment of Fertility: No evidence of carcinogenicity was found in mice or rats following dietary administration of zonisamide for two years at doses of up to 80 mg/kg/day. In mice, this dose is approximately equivalent to the maximum recommended human dose (MRHD) of 400 mg/day on a mg/m² basis. In rats, this dose is 1-2 times the MRHD on a mg/m² basis.

Zonisamide increased mutation frequency in Chinese hamster lung cells in the absence of metabolic activation. Zonisamide was not mutagenic or clastogenic in the Ames test, mouse lymphoma assay, sister chromatid exchange test, and human lymphocyte cytogenetics assay *in vitro*, and the rat bone marrow cytogenetics assay *in vitro*.

Rats treated with zonisamide (20, 60, or 200 mg/kg) before mating and during the initial gestation phase showed signs of reproductive toxicity (decreased corpora lutea, implantations, and live fetuses) at all doses. The low dose in this study is approximately 0.5 times the maximum recommended human dose (MRHD) on a mg/n² basis. The effect of zonisamide on human fertility is unknown.

Pregnancy: Pregnancy Category C (see WARNINGS, Teratogenicity subsection): Zonisamide was teratogenic in mice, rats, and dogs and embryolethal in monkeys when administered during the period of organogenesis. Fetal abnormalities or embryo-fetal deaths occurred in these species at zonisamide dosage and maternal plasma levels similar to or lower than therapeutic levels in humans, indicating that use of this drug in pregnancy entails a significant risk to the fetus. A variety of external, visceral, and skeletal malformations was produced in animals by prenatal exposure to zonisamide. Cardiovascular defects were prominent in both rats and dogs.

Following administration of zonisamide (10, 30, or 60 mg/kg/day) to pregnant dogs during organogenesis, increased incidences of fetal cardiovascular malformations (ventricular

septal defects, cardiomegaly, various valvular and arterial anomalies) were found at doses of 30 mg/kg/day or greater. The low effect dose for malformations produced peak maternal plasma zonisamide levels (25 µg/hl) about 0.5 times the highest plasma levels measured in patients receiving the maximum recommended human dose (MRHD) of 400 mg/day. In dags, cardiovascular malformations were found in approximately 50% of all fetuses exposed to the high dose, which was associated with maternal plasma levels [44 µg/ml] approximately equal to the highest levels measured in humans receiving the MRHD. Incidences of skeletal malformations were also increased at the high dose, and fetal growth retardation and increased frequencies of skeletal variations were seen at all doses in this study. The low dose produced maternal plasma levels [12 µg/ml] about 0.25 times the highest human levels. In cynomolaus monkeys, administration of zonisamide [10 or

In cynomolgus monkeys, administration of zonisamide (10 or 20 mg/kg/day) to pregnant animals during organogenesis resulted in embryo-fetal deaths at both doses. The possibility that these deaths were due to malformations cannot be ruled out. The lowest embryolethal dose in monkeys was associated with peak maternal plasma zonisamide levels [5 µg/ml] approximately 0.1 times the highest levels measured in patients at the MRHD.

at the MRHD.

In a mouse embryo-fetal development study, treatment of pregnant animals with zonisamide (125, 250, or 500 mg/kg/day) during the period of organogenesis resulted in increased incidences of fetal malformations (skeletal and/or craniofacial defects) at all doses tested. The low dose in this study is approximately 1.5 times the MRHD on a mg/m² basis. In rats, in crassed frequencies of malformations (cardiovascular defects) and variations (persistent cords of thymic tissue, decreased skeletal assification) were observed among the offspring of dams treated with zonisamide (20, 60, or 200 mg/kg/day) throughout organogenesis at all doses. The low effect dose is approximately 0.5 times the MRHD on a mg/m² basis.

Perinatal death was increased among the offspring of rats treated with zonisamide {10, 30, or 60 mg/kg/day} from the latter part of gestation up to wearing at the high dose, or approximately 1.4 times the MRHD on a mg/m² basis. The no effect level of 30 mg/kg/day is approximately 0.7 times the MRHD on a mg/m² basis.

There are no adequate and well-controlled studies in pregnant women. ZONEGRAN should be used during pregnancy only if the potential benefit justifies the potential risk to the fetus.

Labor and Delivery: The effect of ZONEGRAN on labor and delivery in humans is not known.

Use in Nursing Mothers: It is not known whether zonisamide is excreted in human milk. Because many drugs are excreted in human milk and because of the potential for serious adverse reactions in nursing infants from zonisamide, a decision should be made whether to discontinue nursing or to discontinue drug, taking into account the importance of the drug to the mother. ZONEGRAN should be used in nursing mothers only if the benefits outweigh the risks.

Pediatric Use: The safety and effectiveness of ZONEGRAN in children under age 16 have not been established. Cases of oligohidrosis and hyperpyrexia have been reported (see WARNINGS, Oligohidrosis and Hyperthermia in Pediatric Patients subsection).

Geriatric Use: Single dose pharmacokinetic parameters are similar in elderly and young healthy volunteers [see CUNI-CAI PHARMACOLOGY, Special Populations subsection in full Prescribing Information). Clinical studies of zonisamide did not include sufficient numbers of subjects aged 65 and over to determine whether they respond differently from younger subjects. 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.

ADVERSE REACTIONS

The most commonly observed adverse events associated with the use of ZONEGRAN in controlled clinical trials that were not seen at an equivalent frequency among placebo-treated patients were somnolence, anorexia, dizziness, headache, nausea, and agitation/irritability.

nausea, and agitation/irritability.

In controlled clinical trials, 12% of patients receiving ZONE-GRAN as adjunctive therapy discontinued due to an adverse event compared to 6% receiving placebo. Approximately 21% of the 1,336 patients with epilepsy who received ZONEGRAN in clinical studies discontinued treatment because of an adverse event. The adverse events most commonly associated with discontinuation were somonlence, fatigue and/or ataxia (6%), anorexia (3%), difficulty concentrating (2%), difficulty with memory, mental slowing, nausea/vomiting (2%), and weight loss (1%). Many of these adverse events were dose-related (see WARNINGS and PRECAUTIONS).

Adverse Event Incidence in Controlled Clinical Trials: Table 3 lists treatment-emergent adverse events that occurred in at least 2% of patients treated with ZONEGRAN in controlled clinical 2% of patients reduced with ZONEGRAN in Continued chitch trials that were numerically more common in the ZONEGRAN group. In these studies, either ZONEGRAN or placebo was added to the patient's current AED therapy. Adverse events were usually mild or moderate in intensity.

were usually mild or moderate in intensity.

The prescriber should be aware that these figures, obtained when ZONEGRAN was added to concurrent AED therapy, cannot be used to predict the frequency of adverse events in the course of usual medical practice when patient characteristics and other factors may differ from those prevailing during clinical studies. Similarly, the cited frequencies cannot be directly compared with figures obtained from other clinical investigations involving different treatments, uses, or investigators. An inspection of these frequencies, however, does provide the prescriber with one basis by which to estimate the relative contribution of drug and non-drug factors to the adverse event incidences in the population studied.

Table 3. Incidence (%) of Treatment-Emergent Adverse Events

Table 3. Incidence (%) of Treatment-Emergent Adverse Events in Placebo-Controlled, Add-On Trials (Events that oc-

curred in at least 2% of ZONEGRAN-treated patients and occurred more frequently in ZONEGRAN-treated than placebo-treated patients)

ZONEGRAN (n=269) PLACEBO (n=230)

ZONEGRAN (n=269) PLACEBO (n=230)

BODY AS A WHOLE Headache (10%/8%), Abdominal Pain (6%/3%), Flu Syndrome (4%/3%) DIGESTIVE Anorexia (13%/6%), Nausea (9%/6%), Diarrhea (5%/2%), Dyspepsia (3%/1%), Constipation (2%/1%), Dry Mouth (2%/1%) HEMATOLOGIS AND LYMPHATIC Ecchymosis (2%/1%) METABOUL AND NUTRITIONAL Weight Loss (3%/2%) MERVOUS SYSTEM Dizziness (13%/7%), Ataxia (6%/1%), Nystagmus (4%/2%), Paresthesia (4%/1%) NEUROPSYCHIATRIC AND COGNITIVE DYSFUNCTION-ALTERED COGNITIVE FUNCTION Confusion (6%/3%), Difficulty Concentrating (6%/2%), Difficulty With Memory (6%/2%), Mental Slowing (4%/2%), Difficulty With Memory (6%/2%), Mental Slowing (4%/2%), Netrousness (2%/1%), Insomnia (6%/3%), Anxiety (3%/2%), Netrousness (2%/1%), Netrousness (2%/2%), Difficulties (2%/2%), Difficulties (2%/2%), Difficulties in Verbal Expression (2%/21%), RESPIRATORY Rhinitis (2%/2%), SIfficulties in Verbal Expression (2%/21%), RESPIRATORY Rhinitis (2%/2%), SIfficulties in Verbal Expression (2%/21%), RESPIRATORY Rhinitis (2%/2%), SIfficulties in Verbal Expression (2%/21%), RESPIRATORY Rhinitis (2%/2%), Stecthal Senses (2%/21%), Difficulties in Verbal Expression (2%/21%), Difficulties

Diplopia (6%/3%), Taste Perversion (2%/0%)

Other Adverse Events Observed During Clinical Trials: ZONEGRAN has been administered to 1,598 individuals during all
clinical trials, only some of which were placebo-controlled.

During these trials, all events were recorded by the investigators using their own terms. To provide a useful estimate of the
proportion of individuals having adverse events, similar events
have been grouped into a smaller number of standardized categories using a modified COSTART dictionary. The frequencies
represent the proportion of the 1,598 individuals exposed to
ZONEGRAN who experienced an event on at least one occasion. All events are included except those already listed in the
previous table or discussed in WARNINGS or PRECAUTIONS,
trivial events, those too general to be informative, and those not
reasonably associated with ZONEGRAN.

Events are further classified within each category and listed in

Events are further classified within each category and listed in order of decreasing frequency as follows: <u>frequent</u> occurring in at least 1:100 patient; <u>infrequent</u> occurring in 1:100 to 1: 1000 patients; <u>rare</u> occurring in fewer than 1:1000 patients.

Body as a Whole: Frequent: Accidental injury, asthenia. Infrequent: Chest pain, flank pain, malaise, allergic reaction, face edema, neck rigidity. Rare: Lupus erythematosus.

Cardiovascular: Infrequent: Palpitation, tachycardia, va insufficiency, hypotension, hypertension, thrombophlebitis, syncope, bradycardia. *Rare:* Atrial fibrillation, heart failure, pulmonary embolus, ventricular extrasystoles.

Digestive: Frequent: Vomiting. Infrequent: Flatulence, gingivitis, gum hyperplasia, gastrilis, gastroenteritis, stomatitis, chole-lithiasis, glossitis, melena, rectal hemorrhage, ulcerative stomatitis, gastro-duodenal ulcer, dysphagia, gum hemorrhage. Rare: Cholangitis, hematemesis, cholecystitis, cholestatic iaundice, colitis, duodenitis, esophagitis, fecal incontinence, mouth ulceration.

Hematologic and Lymphatic: Infrequent: Leukopenia, anemia, immunodeficiency, lymphadenopathy. Rare: Thrombocytopenia, microcytic anemia, petechia.

Metabolic and Nutritional: *Infrequent:* Peripheral edema, weight gain, edema, thirst, dehydration. *Rare:* Hypoglycemia, hyponatremia, lactic dehydrogenase increased, SGOT increased.

Musculoskeletal: Infrequent: Leg cramps, myalgia, myasthenia, arthralaia, arthritis

Nervous System: Frequent: Tremor, convulsion, abnormal goit, hyperesthesia, incoordination. Infrequent: Hypertonia, twitching, abnormal dreams, verligo, libido decreased, neuropathy, hyperkinesia, movement disorder, dysarthria, cerebrovascular accident, hypotonia, peripheral neuritis, parathesia, reflexes increased. Rare: Circumoral paresthesia, dyskinesia, dystonia, encephalopathy, facial paralysis, hypokinesia, hyperesthesia, myoclonus, oculogyric crisis.

Behavioral Abnormalities - Non-Psychosis-Related: Infrequent:

Respiratory: Frequent: Pharyngitis, cough increased. Infrequent: Dyspnea. Rare: Apnea, hemoptysis.

Skin and Appendages: Frequent: Pruritus. Infrequent: Macu-lopapular rash, acne, alopecia, dry skin, sweating, eczema, urticaria, hirsutism, pustular rash, vesiculobullous rash.

Special Senses: Frequent: Amblyopia, tinnitus. Infrequent: Con-junctivitis, parosmia, deafness, visual field defect, glaucoma. Rare: Photophobia, iritis.

Urogenital: Infrequent: Urinary frequency, dysuria, urinary incontinence, hematuria, impotence, urinary retention, urinary urgency, amenorrhea, polyuria, nocturia. *Rare:* Albuminuria, enuresis, bladder pain, bladder calculus, gynecomastia, mastitis, menorrhagia



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