CNS SPECTRUMS[®] The International Journal of Neuropsychiatric Medicine

> Is SAD Lost to SAD? S. Kasper

REVIEW ARTICLES

The Diagnosis, Symptomatology, and Epidemiology of Seasonal Affective Disorder A. Magnusson and T. Partonen

Update on the Biology of Seasonal Affective Disorder C-H Sohn and R.W. Lam

Light Therapy for Seasonal and Nonseasonal Depression: Efficacy, Protocol, Safety, and Side Effects *M. Terman and J.S. Terman*

The Pharmacotherapy of Seasonal Affective Disorder E. Pjrek, D. Winkler, and S. Kasper

CASE REPORT

The Impact of Comorbidity on the Management of Pathological Gambling B. Dell'Osso and E. Hollander

CLINICAL COLUMN

Pearls in Clinical Neuroscience: Suffer the Children: Psychobiology of Early Adversity D.J. Stein, B.A. Harvey, J. Uys, and W. Daniels



Index Medicus/MEDLINE citation: CNS Spectr

When You Treat ADHD....

GF



ADDERALL XR[®] Delivers Efficacy That May Help Patients Realize Their Potential

- Symptom reduction to a level comparable to that of non-ADHD peers'
- Rapid onset (1.5 hours) and 12-hour dose-responsive efficacy for day-long improvement in both academic and social settings^{*2-5}
- · 6 dosage strengths for maximum flexibility
- Generally well tolerated—low discontinuation rates due to adverse events in placebo-controlled trials^{2.4}

*Average mean for all doses tested. *IMS Dataview, May 2005. Please see references and brief summary of prescribing information on adjacent page.

www.ADDERALLXR.com www.ADHDSupport.com

Shire US Inc. ...your ADHD support comj 1-800-828-2088



Reach new heights

Important Safety Information

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. References: I. Ambrosini PJ, Lopez FA, Chandler MC, et al. An open-label community assessment of ADDERALL XR in pediatric ADHD. Poster presented at: 155th Annual Meeting of the American Psychiatric Association; May 22, 2002: Philadelphia, Pa. 2. Data on file, Shire US Inc., 2005. 3. Biederman J, Lopez FA, Boeliner SW, Chandler MC. A randomized, double-blind, placebo-controlled, parallel-group study of SU381 (Adderail XR) in children with attention-deficit/hyperactivity disorder. *Pediatrics*. 2002; 110:258-266. 4. McCracken JT, Biederman J, Greenhill LL, et al. Analog classroom assessment of a once-daily mixed amphetamine formulation, SU381 (ADDERALL XR), in children with ADHD. J Am Acad Child Adolesc Psychiatry. 2003;4:2673-683. 5. Lopez FA, Ambrosini PJ, Chandler MC, et al. ADDERALL XR in pediatric ADHD: quality of life measures from an open-label community assessment trial. Poster presented at: 14th Annual CHADD International Conference; October 17, 2002; Miami Baech, Fla. BRIEF SUMMARY: Consult the full prescribing information for complete product information.

ADDERALL XR® CAPSULES

CII Rx Only

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 EVENTS

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-IV criteria for ADHD (see CLINICAL PHARMACOLOGY), along with extrapolation from the known efficacy of ADDERALL®, the immediate-release formulation of this substance.

CONTRAINDICATIONS

LCM TRAINULATIONS Advanced afteriosclerosis, symptomatic cardiovascular disease, moderate to severe hypertension, hyperthyroidism, known hypersensitivity or idiosyncrasy to the sympa-thomimetic amines, glaucoma. Agitated 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

WARNINGS Psychosis: 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 chronic Sultate is use of stimularts 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. Adderail 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

General: The least amount of ampletamme teasure should be presented of dispensed at one time in order to minimize the possibility of overdosage. Hypertension: Caution is to be exercised in prescribing ampletamines 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 moderation.

Thes: Amphetamines have been reported to exacerbate motor and phonic tics and folurettes synghome. Ineretore, clinical evaluation for tics and Tourette's syndrome in children and their families hould precede use of stimulant medications. International activity of the patient should therefore be cautioned accordingly. **Orug Interactions:** Andipfing agents—Castrointestimal aciditying agents (guarentindine, reserpine, glutamic acid HCI, ascorbic acid etc.) lower absorption of amphetamines. *Urinary aciditying agents*—These agents (ammoni-um childride, sodium acid phosphate, etc.) increase the concentration of the ionized species of the amphetamine molecule, thereby increasing unary excretion. Both groups of agents lower blood levels and efficacy of amphet-amines. *Adrenergic blockers*—Adrenergic blockers are inhibited by amphetamines. *Maininzing agents*— Gastrointestimal alkalinizing agents (sodium bicarbonate, etc.) increase absorption of amphetamines. *Co-adminis-*tration of ADDERALL XR® and gastrointestinal alkalinizing agents, such as antacids, should be avoided. Urinary alkalinizing agents (sodium bicarbonate, etc.) increase the concentration of the non-ionized species of the amphetamine molecule, thereby decreasing urinary excretion. Both groups of agents increase blood levels and therefore potentiate the actions of amphetamines. *Antidepressants, tricyciic*—Amphetamines may enhance the activity of tricycic antidepressants or sympathomimetic agents; d-amphetamine with desiparamine or protriptlyline and possibly other tricycics cause striking and sustained increases into concentration of d-amphetamine in the and ther signs of hypertensive crisis. A variety of toxic neurological effects and malignating the refeted on the real-and possibly other tricycics cause baroptions from adrenergic nerve endings, this can cause headaches and other signs of hypertensive crisis. A variety of toxic neurological effects and malignating the central stimulant effects of amphetamines, and can be used to treat a

produce a synergistic anticonvulsant action. *Proposyphene*—In cases of propoxyphene overdosage, ampheta-mine CNS stimulation is potentiated and fatal convulsions can occur. *Veratum akaloids*—Amphetamines inhibit the hypotensive effect of veratum akaloids. **Drug/aboratory Test Interactions:** Amphetamines can cause a significant elevation in plasma corticosteroid levels. This increase is greatest in the evening. Amphetamines may interfere with urinary steroid determinations. **Carcinogenesis/Mutagenesis and Impairment of Ferlility:** No evidence of carcinogenicity was found in studies in which *Q1*-amphetamine (enantiomer ratio of 1:1) was administered to mice and rats in the deit for 2 years at doses of up to 30 mg/kg/day in male mice, 19 mg/kg/day in temale mice, and 5 mg/kg/day in male and female rats. These doses are approximately 2.4, 1.5, and 0.8 times, respectively, the maximum recommended human dose of 30 mg/kg/ (child) on a mg/m² body surface area basis. Amphetamine, in the enantiomer ratio present in ADDEFALL® (inmediate-release)(d- to 1- ratio of 3:1), was not clastogenic in the mouse bone marrow micronucleus test *in vivo* and was negative when tested in the <u>E</u> cosi component of the Ames test *in vitro*. d1-Amphetamine (1:1 enantiomer ratio) has been reported to produce a positive response in the mouse bone marrow micronucleus test. Amphetamine, in the enantiomer ratio present in ADDEFALL® (inmediate-release) (d- to 1- ratio of 3:1), did not aversely affect fertility or early embryonic development in the rat at doses of up to 20 mg/kg/day (approximately 5 times the maximum recommende human dose of 30 mg/day on a mg/m² body surface area basis). **Pregnancy:** Pregnancy Category C. Amphetamine, in the enantiomer ratio present in ADDEFALL® (d- to 1- ratio of 3:1), had no apparent effects on embryofetal morphological development or survival when orally administred to pregnant rats and rabbits throughout the period of organogenesis at doses of 50 mg/kg/day (approximately 5 times the distowing paremtera

ADVERSE EVENTS

ADVENSE EVENTS 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 6 to 12) were evaluated in two controlled clinical studies, one open-label clinical study, and two single-dose clin-ical 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

sandarder events aregures, in the tables and issuings that follow, costArri 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 insomia) compared to 2.7% (7/259) receiving placebo. The most frequent adverse events associated with discontinuation of ADDERALL XR[®] (controlled and uncontrolled, multiple-dose clinical trials of pediatric patients (n=565) are presented below. Over half of these patients were exposed to ADDERALL XR[®] for 12 months or more.

Adverse event Anorexia (loss of appetite) Insomnia Weight loss Emotional lability Depression

ALL XR[®] (**zs mg, 30 mg CAPSULG** (N=191) were 3.1% (n=6) for nervousness including anxiety and inritability. 2.6% (N=191) were 3.1% (n=6) for nervousness including anxiety and inritability. 2.6% (N=191) were 3.1% (n=6) for nervousness including anxiety and inritability. 2.6% (N=191) were 3.1% (n=6) for nervousness including anxiety and inritability. 2.6% (n=5) for insomnia, 1% (n=2) each for headache, paipitation, and somnolence; and 0.5% (n=1) each for ALT increase, agitation, chest pain, cocaine craving, elevated blood pressure, and weight loss. Adverse events occurring in a controlled trial: Adverse events reported in a 3-week clinical trial of pediatric patients and a 4-week clinical trial in adults treated with ADDERALL XR[®] or placebo are oresented in the tables below.

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presented in the tables below. The prescriber should be aware that these figures cannot be used to predict the incidence of adverse events in the 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 investigators. 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 Higher Incidence Than on Placebo in a 584 Patient Clinical Study

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 Lability	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 an 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%
Urnnenital System	Urinary Tract Infection	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 XR® with a higher incidence than patients receiving placebo in this study: Infection, photosensitivity reaction, constipation, tooth disorder, emotional lability, libido decreased, sormolence, speech disorder, paipitation, witching, dyspina, 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 of cardiomyopathy associated with chronic amphetamine use. Central Nervous System: Psychotic episodes at recom-mended doses, overstimulation, restlessness, dizziness, insomnia, euphoria, dyskinesia, dysphoria, depression, tremor, headache, exacerbation of motor and phonic fics and Tourette's syndrome, selzures, stroke. Gastrointestinai: Dryness of the mouth, unpleasant taste, diarrhea, constitution, other gastrointestinai disturbances. Anorexia and weight loss may occur as undesirable effects. Allergic: Urticaria. Endocrine: Impotence, changes in libido.

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. 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. **OVERODSAGE**

Trom schizophreina. **OVERDOSAGE** Individual patient response to amphetamines varies widely. Toxic symptoms may occur idiosyncratically at low doese. Symptoms: Maniestations of acuts overdosage with amphetamines include restlessness, tremor, hyper-reflexia, rapid respiration, confusion, assaultiveness, hallucinations, panic states, hyperpyrexia and rhabdomyol-ysis. Failgue and depression usually follow the central nervous system stimulation. Cardiovascular effects include arritythmias, hypertension or hypotension and circulatory collapse. Castrointestimal symptoms include nausea, vomiting, diarrhea, and abdominal cramps. Fatal poisoning is usually preceded by convulsions and coma. Teratment: Consult with a Cartified Poison Control Centre for up to date guidance and advice. Management of acute administration of a cathartic and sedation. Experience with hemodialysis or peritoneal dialysis is inadeguate to permit recommendation in this regard. Acidification of the urine increases amphetamine excertion, but is believed to increase risk of acute renal failure if myoglobinuria is present. If acute severe hypertension complicates ampheta-mine overdosage, administration of intravenous phentohamine has been achieved. Chiorpromazine antagonizes the central stimulant effects of amphetamines and can be used to treat amphetamine intoxication. The prolonged release of mixed amphetamine salts from ADDERALL XR® should be considered when treating patients with overdose. Dispense in a hight, 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., Wayne, PA 19087 Made in USA For more information call 1-800-828-2088. or visit www.adderailx.com. ADDERALL XR® are registered 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., Wayne, PA 19087 Made in USA For more information call 1-800





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

% of pediatric patients discontinuing (n=595)

CNS SPECTRUMS

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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residual symptoms sadness low energy anxiety relapse

recurrence

of unresolved depression with EFFEXOR XR12

IMPORTANT TREATMENT CONSIDERATIONS

Suicidality in Children and Adolescents

Antidepressants increased the risk of suicidal thinking and behavior (suicidality) in short-term studies in children and adolescents with Major Depressive Disorder (MDD) and other psychiatric disorders. Anyone considering the use of EFFEXOR XR or any other antidepressant in a child or adolescent must balance this risk with the clinical need. Patients who are started on therapy should be observed closely for clinical worsening, suicidality, or unusual changes in behavior. Families and caregivers should be advised of the need for close observation and communication with the prescriber. EFFEXOR XR is not approved for use in pediatric patients.

EFFEXOR XR is contraindicated in patients taking monoamine oxidase inhibitors (MAOIs). Adult and pediatric patients taking antidepressants can experience worsening of their depression and/or the emergence of suicidality. Patients should be observed closely for clinical worsening and suicidality, especially at the beginning of drug therapy, or at the time of increases or decreases in dose. Anxiety, agitation, panic attacks, insomnia, irritability, hostility, impulsivity, akathisia, hypomania, and mania have been reported and may represent precursors to emerging suicidality. Stopping or modifying therapy should be

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considered especially when symptoms are severe, abrupt in onset, or not part of presenting symptoms. Treatment with venlafaxine is associated with sustained increases in blood pressure (BP) in some patients. Regular BP monitoring is recommended. Abrupt discontinuation or dose reduction has been associated with discontinuation symptoms. Patients should be counseled on possible discontinuation symptoms and monitored while discontinuing the drug; the dose should be tapered gradually. The most common adverse events reported in EFFEXOR XR short-term placebo-controlled depression, generalized anxiety disorder (GAD), and/or social anxiety disorder trials (incidence $\geq 10\%$ and $\geq 2x$ that of placebo) were anorexia, asthenia, constipation, dizziness, dry mouth, ejaculation problems, impotence, insomnia, nausea, nervousness, somnolence, and sweating.

Please see brief summary of Prescribing Information on adjacent pages.

References: 1. Data on file, Wyeth Pharmaceuticals Inc. 2. Effexor XR[®] (venlafaxine HCI) Extended-Release and Effexor Immediate-Release Prescribing Information, Wyeth Pharmaceuticals Inc.





BRIEF SUMMARY. See package insert for full prescribing information

Suicidality in Children and Adolescents

Antidepresents increased the risk of suicidal thinking and behavior (suicidality) in short-term studies in children and adolescents with Major Depressive Disorder (MDD) and other psychiatric disorders. Anyone considering the use of EFFEXOR XR or any other antidepressant in a child or adolescent must belance this risk with the clinical need. Patients who are started on therapy should be observed closely for clinical worsening, suicidality, or unusual changes in behavior. Families and caregivers should be advised of the need for close observation and communication with the prescriber. EFFEXOR XR is not approved for use in pediatric patients. (See Warnings and Precautions: Pediatric Use.)

Pooled analyses of short-term (4 to 16 weeks) placebo-controlled trials of 9 antidepressant drugs (SSRis and others) in children and adolescents with Major Depressive Disorder (MDD), obsessivecompulsive disorder (OCD), or other psychiatric disorders (a total of 24 trials involving over 4,400 patients) have revealed a greater risk of adverse events representing suicidal thinking or behavior (suicidality) during the first few months of treatment in those receiving antidepressants. The average risk of such events in patients receiving antidepressants was 4%, twice the placebo risk of 2%. No suicides occurred in these trials.

Indexide the events of the original rate of average events representing subclast values of a sch events in patients receiving antidepressants was 4%, twice the placebo risk of 2%. No subcles occurred in these traditivity to ventafaxime hydrochiodie or to any excipients in the formulation. Concomitant use in patients taking monomine oxidase inhibitors (MAOI), WARNINGS: Clinical Worsening and Subcle Risk—Patients with major depressive disorder (MDD), both adult and pediatric, may exceptence worsening of their depression and/or the emergence of subcles locations, and this risk may persist until significant remission occurs. There has been a long-standing concern that antidepressants may have a role in inducing, worsening of the psychiatric disorders. It is also unknown whether the subcleading rate advance of subcleading in a schematic subcleading in the emergence of subcleading in a schematic subcleading rate advance of darge transport of dase changes, either increases and the rate of the expectation is down of the expectation is closed at the subcleading in the emergence of subcleading rate advance advance advance of darge transport of dase changes, either increases or decreases. Adults with MDD or comorbid depression in the satting of other psychiatric lines being trade with antidepressants for any molecution should be observed inmital few months of a course of drug therapy, or at times of dase changes, either increases or decreases. Adults with MDD or comorbid depression in the satting of other psychiatric lines exceeding during the initial few months of a course of drug therapy, or at times of dase changes, either increases or decreases. Adults with between the emergence of subcleading values and either the worsening of depression and/or the emergence of subcleading. Consideration should be given to changing the therapeutic regime, including possibly discontinuity the medication, should be given to changing the therapeutic regime, including possibly discontinuits the medication should be given to changing t PRECAUTIONS: General—Discontinuation of Treatment with Effexor XR. Abropt discontinuation or dose reduction of venlafaxine at various doses is associated with new symptoms, the frequency of which increased with increased dose level and longer duration of treatment. Symptoms include agitation, anorexia, anxiety, contision, coordination impaired, diarrhea, dizziness, dry mouth, dysphoric mood, emotional lability, fasciculation, fatigue, headches, hypomania, insominal, intritability, lethargy, nausea, nervousness, nightmares, seizures, sensory disturbances (e.g., paresthesias such as electric shock sensations), somonolence, sweating, timitus, tremor, vertigo, and vomiting. Monitor patients when discontinuing treatment. A gradual reduction in the dose rather than abrupt cessation is recommended. If intolerable symptoms occur following a decrease in the dose or upon discontinuation of threatment, consider resuming the previously prescribed dose. Subsequently, continue decreasing the dose at a more gradual rate. *Insomnia and Nervousness*. Treatment-emergent insomnia and nervousness have been reported. In Phase 3 trials, insomnia led to drug discontinuation in 0.9% of depressed patients. *Changes in Weight: Adult Patients*. In short-term MDD frials, 7% of Effexor XR patients had ≥5% loss of body weight and 0.1% discontinued for weight loss. In 6-month GAD studies, 3% of Effexor XR patients had ≥5% loss of body weight, and 0.3% discontinued for weight loss. In 8-week Subtisel. Changes in Weight Loss alone or in combination with weight loss agents, including phentermine, have not been established. Coadministration of Effexor XR and weight loss agents is not recommended. Effexor XR is not indicated for weight loss alone or in combination with weight loss agents is not recommended. Effexor XR is not indicated for weight loss alone or in combination with weight loss agents is not recommended. Effexor XR is not indicated for weight loss. The setter street week and bright reductive devisites (2.9% of AD patients week SA

syndrome of inappropriate antidiuretic hormone secretion (SIADH) may occur with venlafaxine. Consider this in patients who are volume-depleted, elderly, or taking diuretics. *Mydriasis*: Mydriasis has been reported; monitor patients with raised intraocular pressure or at risk of acute narrow-angle glaucoma). Seturnes: In all premarketing depression trials with Efeors, seizures were reported in 0.3% of venlafaxine patients. Use cautiously in patients with a history of seizures. Discontinue in any patient who develops seizures. Abnormal Bleeding: Abnormal bleeding (most commonly ecchymosis) has been reported. Serum Cholesterol leveration: Clinically relevant increases in serum cholesterol were seen in 5.3% of venlafaxine patients and 0.0% of placebo patients treated for at least 3 months in trials. Consider measurement of serum cholesterol levels during long-term treatment. Use In Patients With Concomitant *Hiness*: Use Effexor XR cautiously in patients with diseases or conditions that could affect hemodynamic responses or metabolism. Venlafaxine has not been reported in clinical studies. Exercise caution in patients whose underlying medical conditions might be compromised by increases in heart rate. In patients with resent with eccent history of MI or unstable impairment or cirrbosis of the liver, the clearances of venlafaxine and its active metabolites were decreased. whose underlying medical conditions might be compromised by increases in heart rate. In patients with renal impairment or cirrhosis of the liver, the clearances of venlafaxine and its active metabolites were decreased, prolonging the elimination half-lives. A lower dose may be necessary; use with caution in such patients. Information for Patients—Prescribers or other health professionals should inform patients, their families, and their caregivers about the benefits and risks associated with treatment with Effexor XR and should counsel them in its appropriate use. A patient *Medication Guide About Using Antidepressants in Children and Teenagers* is available for Effexor XR. The prescriber or health professional should instruct patients, their families, and their caregivers to read the Medication Guide and should assist them in understanding its contents. Patients should be given the opportunity to discuss the contents of the Medication Guide and to obtain answers to any questions they may have. The complete text of the Medication Guide is available at <u>www.effexor.com</u> or the approved prescribing information. Patients should be advised of the following issues and asked to alert should be given the opportunity to discuss the contents of the Medication Guide and to obtain answers to any questions they may have. The complete text of the Medication Guide is available at <u>www.effisoxr.com</u> or the approved prescribing information. Patients should be advised of the following issues and asked to alert their prescriber if these occur while taking Effexor XR. **Clinical Worsening and Suicide** Risk: Patients, their families, and their caregivers should be encouraged to be alert to the emergence of symptoms listed in **WARNINGS: Clinical Worsening and Suicide Risk**, especially those seen early during antidepressant treatment and when the dose is adjusted up or down. Families and caregivers of patients should be advised to observe for the emergence of such symptoms on a day-to-day basis, since changes may be abrupt. Such symptoms should be reported to the patient's prescriber or health professional, especially if they are severe, abrupt in onset, or were not part of the patient's prescriber or health professional, especially if they are severe, including automobiles, until they are reasonably sure that venlafaxine does not adversely affect their abilities. Teil patients to avoid alcohol while taking Effexor XR and to notify their physician 1) if they become pregnant or interd to become pregnant during therapy, or if they are nursing; 2) about other prescription or over-the-counter drugs, including herbal preparations; they are taking or plan to take; 3) if they devole a rash, hives, or related allergic phenomena. Laboratory Tests— No specific Liobaratory tests are recommende. Drug Interactions— Alcohol: A single dose of ethanol had no effect on the pharmacokinetics (FK) of venlafaxine or -desmethylvenlafaxine (ODV), and venlafaxine ddi not exaggerate the psychomotor and psychometric effects induced by dianol. *Climetidime*: Use caution when administering venlafaxine with cimetidine to patients with pre-existing hypertension or hepatic dystunction, and the elderly. *Diazeparn*. A si Venlataxine is not highly bound to plasma proteins; coadministration of Effexor XR with a highly protein-bound drug should not cause increased free concentrations of the other drug. *Drugs that inhibit Cytochrome P450 Isoenzymes*: CYP2D6 inhibitors: Venlataxine is metabolized to its active metabolite, ODV, by CYP2D6. Drugs inhibiting this isoenzyme have the potential to increase plasma concentrations of venlataxine and decrease concentrations of ODV. No dosage adjustment is required when venlataxine is coadministered with a CYP2D6 inhibitor. Concomitant use of venlataxine with drug treatment(s) that potentially inhibits both CYP2D6 and CYP3A4, the primary metabolizing enzymes for venlataxine, is an to been studied. Use caution if therapy includes venlataxine and any agent(s) that produces simultaneous inhibition of these two enzyme systems. *Drugs Metabolized by Cytochrome P460 Isoenzymes*. Venlataxine is a relatively weak inhibitor of CYP2D6. Hentatxine did not inhibit CYP1A2 and CYP3A4, CYP2O9 (in vitro), or CYP2C19. *Impramine*: Venlataxine did not affect the PK of venlataxine. The 2-OH-designamine AUCs increased by 2-54-5.5 fold. Impramine did not affect the PK of venlataxine in the 2-OH-designation experime AUCs. *Cynas and Gyna*, increase in risperidone bit as active metabolite, 9-hydroxyrisperidone, resulting in a -32% increase in risperidone plus 9-hydroxyrisperidone). CYP3A4: Venlataxine did not inhibit CYP3A4 in vitro and in vico. *Indinavir* in a study of 9 healthy volunteners, venlataxine administration resulted in 28% decrease in the AUC of a single close of indinavir and a 36% decrease in indinavir Cima. Indinavir did not affect the PK of venlataxine did not inhibit CYP2C9: Venlataxine did not inhibit CYP2C9 venlataxine did not inhibit CYP2C9 in vitro. In vivo. CYP2C919: Venlataxine did not inhibit CYP2C9 in vitro. In vivo. Venlataxine did not inhibit CYP2C9 in vitro. In vivo. Venlataxine did not inhibit CYP2C9 in vitro. In vivo. CYP2C919: Venlataxine did not inhibit CYP2C9 in vitr begin burking pressions and pression of the second second

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CNS SPECTRUMS The International Journal of Neuropsychiatric Medicine

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672 The quiz on seasonal affective disorder is CME-accredited by Mount Sinai School of Medicine for 3.0 credit hours.

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Volume 10 - Number 8

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SEROQUEL is indicated for the treatment of acute manic episodes associated with bipolar I disorder, as either monotherapy or adjunct therapy with lithium or divalproex, 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. A rare condition referred to as neuroleptic malignant syndrome has been reported with this class of medications, including SEROQUEL.

Hyperglycemia, in some cases extreme and associated with ketoacidosis, hyperosmolar coma, or death, has been reported in patients treated with atypical antipsychotics, including SEROQUEL. Patients starting treatment with atypical antipsychotics who have or are at risk for diabetes should undergo fasting blood glucose testing at the beginning of and during treatment. Patients who develop symptoms of hyperglycemia should also undergo fasting blood glucose testing.

Precautions include the risk of seizures, orthostatic hypotension, and cataract development.

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

© 2005 AstraZeneca Pharmaceuticals LP. All rights reserved. SEROQUEL is a registered trademark of the AstraZeneca group of companies. (/doi.org/10.1017/S1692525000(13507 Published online by Cambridge University Press BRIEF SUMMARY of Prescribing Information-Before prescribing, please consult complete SEROQUEL® (quetiapine furnarate) Tablets Prescribing Informati

INDICATIONS AND USAGE: Bipolar Mania: SEROQUEL is indicated for the treatment of acute manic INDICATIONS AND USAGE: Bipolar Mania: SERQUUEL is indicated for the treatment of acute mani-episodes associated with bipolar i losiondr, as either monotherapy or adjunct therapy to tilhium c divalorex. The efficacy of SERQUUEL in acute bipolar mania was established in two 12-we montherapy traits and nea 3-week adjunct therapy that of bipolar plates initially hospitaized for up to 7 days for acute mania. Effectiveness has not been systematically evaluate in clinical traits to more than 12 weeks in monotherapy or adjunct therapy. Therefore, the physician will elects to use SERDOUEL for extended periods should periodically re-avaluate the long-term risks an election to use SERDOUEL for extended periods should periodically re-avaluate the long-term risks and controller that a of chapter mice lighters. Therefore therapients: SERDOUEL in non-term (3-week controller that as of chaptering tiltients. Therefore therapients of SERDOUEL in non-term use, that is for more than 6 weeks, has not been systematically evaluated in controlled trais. Therefore, the physical is use SERDOUEL for extended periods should periodically re-valuate the long-term is should be solvebornein lighters. Therefore theremess of SERDOUEL in non-term is week in some than 6 weeks, has not been systematically evaluated in controlled trais. Therefore, the physical is and voletist to use SERDOUEL in centende period should periodically re-valuate the long-term usefulness of the drug for the individual patient.

CONTRAINDICATIONS: SEROQUEL is contraindicated in individuals with a known hypersensitivity to

sakuness of the drug for the individual patient. **CONTRAINDICATIONS:** SEROUUEL is contraindicated in individuals with a known hypersensitivity to the medication or any of its ingredients. **WARNIOS:** Neuroispite Malignent Synchrom (MNS): A potentially fatal symptom complex some-mines refere to as MISA has been reported in association with administration of antipsycholic drugs, induling SEROUUEL. Bare cases of INIS have been reported with SEROUEL. Clinical manifesta-ment of the analysis of the synchrome (MNS): A potentially fatal symptom complex some-instability. See full Prescription information for more information on the manifestations, diagnosis and management of NNS. If a patient requires antipsycholic drug trainment after recovery from MNS. If the patient requires antipsycholic drug trainment after recovery from MNS. Its patient requires antipsycholic drug trainment after recover from MNS. If the patient requires antipsycholic drug trainment after recovery from MNS. If the patient requires antipsycholic drug trainment after recovery from MNS. If the patient agriculture trained with the patients are likely to develop the synthesis. A syndrome of potentially interestible, myolumiany, dyskinetic movements may develop in patients treated with antipsycholic drugs. Although the prevalence of the syndrome apparts to the highest among the ellery's products differ in their potential to cause tardwor dyskinesis is unknown. The risk of developing trained systhmest and the likelihout that will baccome inversible are believed to interase as the duration of treatment which accumulative dose of antipsycholic trainers is unbiggeduit trainers is sittly drawed dyskinese and the likelihout that will baccome intervisible accumoly, after trainers is sittly drawed dyskinese and the likelihout the distly completely. If antipsycholic trainers that be applete to trainers the trainers and the systhmest are not variable or sporoniae. There is no known treatment that is most likely to minimize the courserve of

Precoursons: some callents required continuation of anti-diabetic treatment despite discontinuation of the suspect drug. Precoursons: the suspect drug. Precoursons: the suspect drug and the suspect drug and the suspect drug and the suspect drug. Precoursons: the suspect drug and the suspective drug and the suspect drug and the suspect drug. In some patients: synchose and the suspective drug and the suspective drug and the suspect drug and the suspect drug. In some patients with norw cardiovascular disease (history of mycoratial infarction or ischemic heart disease, heart later or conduction anomalities), caredovascular disease or conditions which would predispose patients to thy potension (dehydration, hypovolenia and survece) may be minimized by limiting the initial does to 25 mp bit. If hypotension course during titration to the larget does, a return to the previous dises to 25 mp bit. If hypotension course during titration to the larget does, a return to the previous dises to 25 mp bit. If hypotension course during titration to the larget does, a return to the previous dises to 25 mp bit. If hypotension course during titration to the larget does, are turn to the previous dises to 25 mp bit. If hypotension course during titration to the larget does, are turn to the previous dises to 25 mp bit. If hypotension course during titration to the larget does, are turn to the previous dises to 25 mp bit. If hypotension course during titration to the larget does, are turn to the previous dises to 25 mp bit. If hypotension does course during titration to the larget does, are turn to the previous dises to 25 mp bit. If hypotension does course during the scale and the distribution developments, the associated and the distribution development does and 0.7% (49227) on active associated with a strenge to 25 (19607) to placebo and 0.7% (49227) on active course during the does are obtained development during the does are obtained development of the during down of the during the does are obtain development during down approximately 6% for SERDOUEL compared to 1% for placebo, lin acuté bipolar mania trials, the pro-protinos of patients with transmissional selevations of 3.5 times the upper limits of the normal reference range in a pool of 3- to 12-week placebo-controlled trials were approximately 1% for both SEROUEL and placebo. These headic enzyme elevations usually occurred within the first 3 week of drug trast-ment and promptly network to pre-study levels with orgoing trastmert with SEROUEL. Pointuit of placebo platents trastate dwth SEROUEL especially during the 5-5 diverse event reported in placebo. These headic enzymert. Somolence was a commonly reported adverse event reported in placebo. These headic enzymert. Somolence was a commonly reported adverse event reported in placebo platents. In acute bipolar mania trials using SEROUEL as monotherapy, somolence was reported in 16% of patients on SEROUEL compared to 4% of placebo patients. In acute bipolar main traits using SEROUEL aspecially during the 5-5 diverse platents. In acute bipolar main traits using SEROUEL adverse adjunct therapy, somolence was reported in 34% of placebo patients of the second state of the second state bipolar merit. Thinking, SEROUEL adverse platents should be calined adverse second the requiring merit al alertness, such as operating a motor vehicle (including automobiles) or operating haardous strotoclic compared to 5% of placebo patients. Since SEROUEL has the potential to timpair updy-merit. Thinking a causaria relations for use of SEROUEL has the networking requiring merital alertness. Such as operating a motor vehicle (including automobiles) or operating haardous with alpha-admengic blocking effects have been reported to fudue origins, and it is possible that strotoclicin. White a causari relations for use of SEROUEL has not been established, other drugs with alpha-admengic blocking effects have been reported to thuice prigram, and it is possible that reque core boty temporature has been attitude to antiskychobic agenty, exposure jo active

Bertoutes - queues a common case or monochigy and montality in electry patients in particular those with advanced Athehemer 6 demandia. SEPOULE, and other antipsycholic drogs should be used auticulary in patients at risks to sporting the provide structure in the patients should account the patient in papier document and burgery provide and the patients should account the patients of the patient structure in order to reduce the document. September 2000 and the patient structure in order to reduce the document and the patient structure in the patients and the patients structure the patients with Case September 2000 and the patient structure in order to reduce the document and the patient structure in the patients in the patients and the patients and structure in the patient structure in the patients and the patients for whom a structure in the patient structure (concurrent) field be used when its structure is the structure in the structu

plasma clearance of SEROQUEL was reduced by 30% to 50% in elderly patients when compared to younger patients. ADVERSE REACTIONS: The information below is derived from a clinical trial database for SEROQUEL constitution of over 3000 patients. Of these approximately 3000 subjects, approximately 2700 (2300 m schophrenia and 45 in acute block mana) were patients who patrolicaled in multiple toose effect trearess traits, and their experience corresponded to approximately 514.3 patient/years. Refer to the UII Preschoing (Information for defails of adverse event data collection. Adverse Findinger, Buserved in Short-Term, Pacebo-Cantrolled trials: Baptar Manel: Overall, discontinualiton due to adverse events were 5.7 % for SEROULEL vs. 5.1% for piacebo in monotherapy and 3.8% for SEROULEL vs. 5.9% for placebo classified in during the other schematic schema (Serong Hang) and a schema events (4%, for SEROULEL vs. 3% for classed) in a gool of com-trolled trials. However, discontinuations due to sommolence and hypotension were considered to be drug related (see PHECAUTIONS): Sommolence 0.8% vs.0% for placebo and Hypotension 0.4% vs. 5% for placebo. Adverse Events (A% for SEROULEL vs. 3% for placebo and Hypotension 0.4% vs. 5% for placebo. Term, Placebo-Controlled Trials: Repair Manel with SEROULEL vs. 3% so the schema to counter during acute therapy of with SEROULEL (dases ranging from 75 to 30 mg/day) where the incidence of this Stepplare with SEROULEL (dases ranging from 75 to 30 mg/day) where the incidence in platebo-trated with SEROULEL (dases ranging from 75 to 30 mg/day) where the incidence and Hypotennia and Bipplare Mania (montherapy): Body as a Whole: Headache, Pan, Asthena, Abdomian Pain, Back Pain, Fever, Farallowascular: Ladycardar, Parkal Hypotension (Disetive: Dry Mouth, Consignation, Vormiling, Dyspessia, Gastroenterling, Gamma Glutarny, Transpeptidase increased; Metabolic and Matribaei: Weight Gain, SCP1 Increased, SGOT Increased; Nervus: Agitation, Sommolence, Dizmers, Anveicy:

SEROQUEL® (quetiapine furnarate) Tablets

BEHOULLE* (quetapine turnarize) (abeles)
Resprincer, Francy, Bis, Phintis, Stan Adponéngez, Rasis, Special Senser, Ambyopai, In these studies, the most commonly observed at view of SROULLE, at sets two thind of placebover somolene (18%), dozines (11%), dozines in the studies of the standard (%), weight gain (S3), dozines (11%), dozines (11

and Steven Johnson syndrome (SLS). DBUG ABUSE AND DEPENDENCE: controlled Substance Class: SEROOUEL is not a controlled sub-stance. Physical and Psychologic dependence: SEROOUEL has not been systematically studed in animals or humans, for this potiential for abuse, tollenance or physical dependence. While the clinical trials did not reveal any tendency for any drug-seeking behavior, these observations were not systematic and its not possible to predict on the basis of this limited experiment die texteril to a clinical control of the systematic and and a substance on survival. Consequently, patients should be evaluated carefully for a history of drug abuse; and such patients should be observed close-ly for signs of misuse or abuse of SEROOUEL, s.g., development of tolerance, increases in dose, drug-seking behavior. hehavio

If you can be appendix of the second Indicatory Contexe and the extension with appropriate instances solution in more than the anomalous and the sympathonimite's agents (epineprint) and dopamile solution to be used, incore beta simulation may worsen hypotension in the setting of quetaprine-induced alpha blockade). In cases of severe extravyoratidal symptoms, antichollinergic medication should be administered. Close medical super-vision and monitoring should continue until the patient recovers.

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