CNS SPECTRUMS[®]

ORIGINAL RESEARCH

Mixed Amphetamine Salts Extended-Release in the Treatment of Adult ADHD: A Randomized, Controlled Trial

R.H. Weisler, J. Biederman, T.J. Spencer, T.E. Wilens, S.V. Faraone, A.K. Chrisman, S.C. Read, and S.J. Tulloch

REVIEW ARTICLES

Longitudinal Studies of PTSD: Overview of Findings and Methods

T. Peleg and A.Y. Shalev

Memory of the Traumatic Event as a Risk Factor for the Development of Posttraumatic Stress Disorder: Lessons from the Study of Traumatic Brain Injury

S. Gil, Y. Caspi, I. Ben-Ari, and E. Klein

Sleep Disturbances in the Aftermath of Trauma and Posttraumatic Stress Disorder

T.A. Mellman and M.M.S. Hipolito

Injury Increases the Risk for PTSD: An Examination of Potential Neurobiological and Psychological Mediators

D. Koren, D. Hemel, and E. Klein

PEARLS IN CLINICAL NEUROSCIENCE

The Cognitive-Affective Neuroscience of the Unconscious

D.J. Stein, M. Solms, and J. van Honk

Index Medicus/MEDLINE citation: CNS Spectr

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STICK IT TO ABHD.

15-mg patch (actual size)



Now Available The First and Only ADHD Patch[™]

The history of safety and efficacy of methylphenidate' in a patch

Reduced core ADHD symptoms for 12 hours when worn for the recommended 9 hours²⁴

-Effects were seen at the first time point measured—2 hours²⁻⁴

May be removed earlier than 9 hours⁴ –If a shorter duration of effect is desired⁴ –If late-day side effects appear⁴



Important Safety Information: CNS Stimulants: Patients with structural cardiac abnormalities or other serious cardiac problems should generally not be treated with stimulants. Physicians should take a careful patient history, including family history, and physical exam, to assess the presence of cardiac disease. Patients who report symptoms of cardiac disease such as exertional chest pain and unexplained syncope should be promptly evaluated. Use with caution in patients whose underlying medical condition might be affected by increases in blood pressure or heart rate.

Daytrana: Patients with allergies to methylphenidate or other ingredients in Daytrana should not receive Daytrana. Skin irritation or contact sensitization may occur. Patients should avoid applying any external heat to the Daytrana patch.

Common adverse events reported by patients who received Daytrana in clinical trials were decreased appetite, insomnia, nausea, vomiting, decreased weight, tics, affect lability, and anorexia, consistent with adverse events commonly associated with the use of methylphenidate.

Methylphenidate: Chronic abuse of methylphenidate can lead to marked tolerance and psychological dependence. Careful supervision following withdrawal from abuse is warranted, as severe depression may occur. Methylphenidate should not be used in patients with marked agitation; glaucoma; tics, diagnosis or a family history of Tourette's syndrome; or current/recent use of monoamine oxidase inhibitors (MAOIs). Frank psychotic episodes, new psychosis, mania, aggression, growth suppression, and visual disturbances have been associated with the use of stimulants and discontinuation of treatment may be appropriate. Use with caution in patients with a history of: psychosis; seizures/EEG abnormalities; bipolar disorder; depression; drug dependence or alcoholism. Hematologic and growth monitoring are advised during prolonged therapy.

Please see Brief Summary of Prescribing Information on adjacent page.

References: 1. Elia J. Attention deficit/hyperactivity disorder: pharmacotherapy. Psychiatry. 2005;2:27-35. 2. Data on file, Shire US Inc, 2006. 3. McGough JJ, Wigal SB, Abikoff H, et al. A randomized, double-blind, placebo-controlled, laboratory classroom assessment of methylphenidate transdermal system in children with ADHD. J Atten Disord. 2006;9:476-485. 4. Daytrana [package insert]. Wayne, Pa: Shire US Inc; 2006.

Daytrana[™] is a trademark of Shire Pharmaceuticals Ireland Limited. **www.Daytrana.com** Shire US Inc. ...your ADHD Support Company[™] 1-800-828-2088 ©2006 Shire US Inc., Wayne, Pennsylvania 19087 D338 06/06 C



Deytrana[™] (methylphenidate transfermal system) INDICATION AND USAGE Attennion Deitell Hyperactivity Disorder (ADHD): Daytrana[™] (methylphenidate transfermal system) is indicated for the treatment of Attention Deficit Hyperactivity Disorder (ADHD) and is available in 10, 15, 20, and 30 mg dosing strengths. The effcacy of Daytrana[™] was established in two controlled chincuit triais in chindren with ADHD. Special Diegnostic Considerations: Specific etiology of this syndrome is unknown, and there is no single diagnostic test. Adequate diagnosis requires the use not only of medical but of special psychological, educational, and social resources. Learning may or may not be impaired. The diagnosis must be based upon a complete history and evaluation of the child and not solely on the presence of the required number of DSM-1/1-TR® characteristics. Need for Comprehensive Treatment Program: Daytrana[™] is indicated as an integral part of a total treatment program to any include other measure (psychological, ductational, social) for patients with this syndrome. Drug treatment may not be indicated for all children with this syndrome. Stimulants are not intended for use in the child who exhibits symptoms secondary to environmental factors and/or other primary psychiatric disorders, including psychosis. Appropriate aducational placement is essential and psychosocial intervention is often helpful. When remedial measures alone are isufficient, the effectiveness of Daytrana[™] for long-term use, i.e., for more than 7 weeks, has not been systematically evaluated in controller traits. The physician who elects to use Daytrana[™] for extended periods should periodically re-evaluate the long-term uselviness of Daytrana[™] for long-term use, i.e., for more than 7 weeks, has not been systematically evaluated in controller traits. The physician who elects to use Daytrana[™] for extended periods should periodically re-evaluate the long-term uselviness of Daytrana[™] for long-term use, i.e., and and adjuttion, sinc

Agitation: DaytranaTM is contraindicated in patients with marked anxiety, tension, and agitation, since the drug may aggravate these

Apitation: Daytrana™ is contraindicated in patients with marked anxiety, tension, and agitation, since the drug may aggravate these symptoms. Hypersensitivity to Methylphenidate: Daytrana™ is contraindicated in patients known to be hypersensitive to methylphenidate or other components of the product (polyester/ethylene vinyl acetate laminate film backing, acrylic adhesive, silcone adhesive, and fluoropolymer-coated polyester). Glaucoma: Daytrana™ is contraindicated in patients with glaucoma. Tise: Daytrana™ is contraindicated in patients with motor lics or with a family history or diagnosis of Tourette's syndrome (see ADVERSE REACTIONS). Monoamine Oxidase inhibitors: Daytrana™ is contraindicated during treatment with monoamine oxidase inhibitor (hypertensive crises may result).

may result). WARNINGS

WARNINGS Serious Cardiovascular Events Sudden Death and Pre-existing Structural Cardiac Abnormalities or Other Serious Heart Problems Children and Adolescents Sudden death has been reported in association with CNS stimulant treatment at usual doses in children and adolescents with structural cardiac abnormalities or other serious heart problems. Although some serious heart problems alone carry an increased risk of sudden death, stimulant products generally should not be used in children or adolescents with known serious structural cardiac abnormalities, cardiomyopathy, serious heart rhythm abnormalities, or other serious cardiac problems that may place them at increased vulnerability to the sympathomistic effects of a stimulant drug. Adults

problems that may place them at increased vulnerability to the sympathomimetic effects of a stimulant drug. Adults Sudden deaths, stroke, and myocardial infarction have been reported in adults taking stimulant drugs at usual doses for ADHD. Athough the role of stimulants in these adult cases is also unknown, adults have a grater likelihood than children of having serious structural cardiac abnormalities, cardiomyopathy, serious heart rhythm abnormalities, cornoray artery disease, or other serious cardiac problems. Adults with such abnormalities should also generally not be trated with stimulant drugs. *Hypertension and Other Cardiovscular Conditions* Simulant metications cause a modest increase in average blood pressure (about 2-4 mmHq) and average heart rate (about 3-6 bpm) (see **ADVERSE REACTIONS**), and individuals may have larger increases. While the mean changes alone would not be sure. Caution is indicated in treating patients should be monitored for larger changes in heart rate (aboud pre-sure. Caution is indicated in treating patients whose underlying medical conditions might be compromised by increases in blood pressure or heart rate, e.g., those with pre-existing hypertension, heart failure, recent myocardial infarction, or **Assessing Cardiovscular Situus in Patients Being Treated With Stimulant Medications**

biod pressure or near rate, e.g., those with pre-existing hypertension, heart tailure, recent myocardial infarction, or ventricular arrhythmia. Assessing Cardiovascular Status in Patients Being Treated With Stimulant medications should have a careful his-forly including assessment for a family history of sudden death or ventricular arrhythma) and physical exam to assess for the presence of cardiac disease, and should receive further cardiac evaluation if findings suggest such disease (e.g., electro-cardiogram and echocardiogram). Patients who develop symptoms such as exertional chest pain, unexplained syncope, or other symptoms suggestive of cardiac disease during stimulant treatment should undergo a prompt cardiac evaluation. **Contact Sensitizations:** Use of Daytrana¹¹ may lead to contact sensitization. Daytrana⁴¹ should be discontinued if contact sensitization is suspective. Chrisma is commonly seen with use of Daytrana⁴¹ and is not by itself an indication of sensitization is suspective of significantly improve within A hours or spreased beyond the patch site. Diagnosis of allergic contact dermatitis should be suspected if erythema is accompanied by evidence of a more intense local reaction (edema, paules, vesiciations of systemic sensitization or with any induce a farter-up of providos are taken with other rules, e.g., and Manifestations of systemic sensitization or with any induce a farter-up of providos dematition of notic positive patch-text sites, or generalized sin eruptions in previously unaffected skin. Other systemic reactions may induce headebe, fore, making, arthraiga, darrhea, or vonting.

periarized skin eruptions in previously unaffected skin. Other systemic reactions may include neauexire, invert, mease, aumeyee, diarrhea, or vomiting. Patientis who develop contact sensitization to Daytrana™ and require oral treatment with methylphenidate should be initiated on oral medication under close medical supervision. It is possible that some patients sensitized to methylphenidate by exposure to Daytrana™ may not be able to take methylphenidate in any form. A study designed to provoke skin sensitization revealed a signal for Daytrana™ to be an irritant and also a contact sensitizer. This study involved an induction phase consisting of continuous exposure to the same skin site for 3 weaks, followed by a 2 weak rest period, and then challenge/rechallenge. Under conditions of the study. Daytrana™ was more initiating than both the placebo patch control and the negative control (saline). Of 133 subjects who participated in the challenge phases of the sensitization or close sensitization is when Daytrana™ as prescribed, alternating assessed for sensitization in the clinical effectiveness studies, it is unknown what the true incidence of particular systemic assess of contact sensitization is when Daytrana™ is used as directed. **Perchaints Adverse Events**

The first microence of semilarization is when bayrana to down as one care. Pre-Existing Psychosis Administration of stimulants may exacerbate symptoms of behavior disturbance and thought disorder in patients with a pre-

Administration of stimulants may exacerbate symptoms of behavior disturbance and thought disorder in patients with a pre-existing psycholic disorder. Biplair Illness Particular care should be taken in using stimulants to treat ADHD in patients with comorbid bipolar disorder because of com-erm for possible induction of a mixed/manic episode in such patients. Prior to initiating treatment with a stimulant, patients with comorbid depressive symptoms should be adequately screened to determine if they are at risk for bipolar disorder, such screening should include a detailed psychiatric history, including a family history of sucide, bipolar disorder, such Emergence of New Psycholic or mains Symptoms, e.g., hallucinations, deusional thinking, or mania in children and adoles-cents without a prior history of psychotic illness or mania can be caused by stimulants at usual doses. If such symptoms occur, consideration should be given to a possible causal role of the stimulant, and discontinuation of treatment may be appropriate. In a pooled analysis of multiple short term, placebo-controlled studies, such symptoms occurred in about 0.1% (4 patients with events out 0.3 428 exposed to methypheniate ar omphetamine for several weeks at usual doses) of stim-ulant-treated patients compared to 0 in placebo-treated patients. Aggression

(4) patients with events Out of 3,452 exposed to meruphyterine or meruphyterine or sector and a sector of the s

Drug Dependence Daytrant¹⁴ should be given cautiously to patients with a history of drug dependence or alcoholism. Chronic abusive use can lead to market tolerance and psychological dependence with varying degrees of abnormal behavior. Frank psychotic episodes can occur, especially with parenteral abuse. Careful supervision is required during withdrawal from abusive use, since severe depression may occur. Withdrawal following chronic therapeutic use may unmask symptoms of the underlying disorder that may require follow-up.

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CII Rx Only

Methylphenidate may decrease the effectiveness of drugs used to treat hypertension. Human pharmacologic studies have shown that methylphenidate may inhibit the metabolism of coumarin anticoagulants, anticonvulsants (e.g., bienobarbital, phenytoin, prindione), and some tricyclic drugs (e.g., impiramine, clomipramine, desipramine) and selective seriotinn reuptake inhibitors. Downward dose adjustemats of these drugs may be required when given concomitantly with methylphenidate. It may be necessary to adjust the dosage and monitor plasma drug concentrations (or, in the case of coumarin, cougulation times), when initiating or discontinuing methylphenidate. Serious adverse events have been reported in conconitant use of methylphenidate with clonidine, atthough no causality for the combination has been estabilised. The safety of using methylphenidate in combination with clonidine atthough no causality for the alpha-2-agonists has not been systematically evaluated. **Carcinogenecits, Mutagenessis, and Impairment of Fertility:** Carcinogenicity studies of transdermal methylphenidate have not been performed. In a lifetime carcinogenicity study of oral methylphenidate carried out in BGC3F Imises, methylphenidate is approximately 60 mg/kg/day, Hepatoblastoma is a relatively rare rodent malignant tumor type. There was no increase in hopatocluluar adomonas and, in males only, an increase in hepatic tumors. The mouse strain used is sensitive to the development of hepatic tumors and the significance of these results to humans is unknown. Toally administered methylphenidate as in the transgenic mouse strain p53°, which is sensitive to genotoxic carcinogenicity study in the transgenic mouse strain p53°, which is sensitive to penotoxic carcinogens, there was no evidence of carcinogenicity. In this study, male and female mice were fiel diets containing the same concentration of methylphenidate as in the lifetime carcinogenicity study; the high-dose groups were exposed to 60 to 74 mg/kg/day of methylphenidate.

Thanster overy cells. Methylophenidate did not impair fertility in male or female mice that were fed diets containing the drug in an 18-week Continuous Breeding study. The study was conducted at doses up to 160 mg/kg/day.

Metrylphenidate did not impair fertility in male of temate mice that were ted diels containing the drug in an 18-week Continuous Breeding study. The study was conducted at doess up to 160 mg/kg/day. **Pregnancy Pregnancy Category C:** Animal reproduction studies with transfermal methylphenidate have not been performed. In a study in which oral methylphenidate was given to pregnant rabbits during the period of organogenesis at doese up to 200 mg/kg/day. Teratogenic effects were seen, although an increase in the incidence of a variation, diation of the lateral vertrickes, was seen at 200 mg/kg/day, this does also produed matemal toxicity. A previously conducted study in rabbits showed teratogenic effects of or granogenesis at doese up to 200 mg/kg/day, no teratogenic effects were seen athough a slight delay in fatal skeletal solitation of the lateral vertices was seen at 000 mg/kg/day, and above, these doess caused some matemal toxicity. In a study in which oral methylphenidate was given to trais throughout pregnancy and lactation at doese up to 60 mg/kg/day, dispiration were accessed at 40 mg/kg/day and above, these doess caused some matemal toxicity. Adeguate and well-controlled studies in pregnant women have not been conducted. Deytrana^m should be used during pregnancy only if the potential benefit usafits the potentian risk to the fetus. **Namile Mothers:** Its not known whether methylphenidate is socreted in human milk. Because many drugs are excreted in human **Marking Mothers:** Its not known whether methylphenidate is socreted in human milk. Because many drugs are excreted in human **Marking Mothers:** Its not known whether methylphenidate is socreted in human milk. Because many drugs are excreted in human fedest of methylphenidate in children nave not been vell established (see **WARNINGS)**. In a study conducted in young rats, methylphenidate was administered orality at doess of up to 100 mg/kg/day for 9 weeks, starting early in the postnalal period (Postnata) Day 71 and continuing through sexual

5 mg/kg/day. The clinical significance of the long-term behavioral effects observed in rats is unknown. **DVFRSE FLACTIONS** The pre-marketing clinical development program for DaytranTM included exposures in a total of 1,155 participants in clinical trials (755 perticing clinical and 400 healthy dult subjects). These participants received DaytranaTM in patch sizes ranging from 6.25 cm² to 50 cm³. The 758 pediatric patients (age 6 to 16 years) were evaluated in 9 controlled clinical studies, 2 open-table clinical studies, and 4 clinical pharmacology studies. Adverse reactions were assessed by collecting adverse events data the results of physical examinations, vital signs, weights, laboratory analyses, and ECGs. **Adverse Findings in Clinical Trials With DaytranaTM Adverse Findings in Clinical Trials With DaytranaTM Adverse Findings in Clinical Trials With DaytranaTM Adverse Events Sacotated With Discontinuouslion of Treatment:** In a 7-wesk double-blind, parallel-group, placebo-controlled study in children with ADHD conducted in the outpatient setting, 7.1% (7/98) of patients treated with DaytranaTM (sicontinued use o adverse events compared with 1.2% (1765) receiving placebo. The reasons for discontinuation among the patients treated with DaytranaTM were application site reptients application site reaction, corrlusional state, crying, its, headaches, irritability, infectious mononucleosis, and viral infection. **Adverse Events Coursing at an incidence of 5% or More Among Patients Treated With DaytranaTM table 1 - unergent study in children with ADHD conducted in the outpatient is atmose for discouble-blind, parallel-group, placebo-controlled study in children with ADHD conducted in the outpatient setting. TRE 5 + Mach Coursenb Hoarded Treatment Expensed Hoardee Mith DaytranaTM is a dermal irritanti.**

	lost Commonly Reporte 6 and 2x Placebo) in a 7			clinical efficacy study had minimal to defin
) of Subjects	erythema. This erythema generally caused no
			iverse Events	minimal discomfort and did not usually interfe with therapy or result in discontinuation fro
Adverse Eve	ent	Daytrana™	Placebo (N = 85)	treatment. If erythema, edema, and/or papul
Mumber of I	Cubianta Mith > 1 Adva	(N = 98)		 do not resolve or significantly reduce within
	Subjects With ≥ 1 Adve			hours after patch removal, further evaluati should be sought. Erythema is not by itself
	usea mitino	12 (12)	2 (2)	indication of contact sensitization. Howev
	sopharyngitis	5 (5)	<u>4 (5)</u> 2 (2)	 sensitization should be considered if eryther
	sopharyngnis eight decreased	9 (9)	0 (0)	 is accompanied by edema, papules, vesicles,
	orexia	5 (5)	1 (1)	 other evidence of more intense local reaction Diagnosis of allergic contact dermatitis should be addressed on the second se
	creased appetite	25 (26)	4 (5)	be corroborated by appropriate diagnostic te
	ect lability*	6 (6)	0 (0)	ing (see WARNINGS - Contact Sensitization
Ins	somnia	13 (13)	4 (5)	Adverse Events With the Long-Term Use
Tic		7 (7)	0 (0)	Daytrena™: In a long-term open-label study - Lup to 40-month duration in 191 children w
Na	sal congestion	6 (6)	1 (1)	ADHD, the most frequently reported treatment
tionally sens mittent emo id headache		onal instability, emotional instability, emotional total of 45 (24%) su	onal lability, and inter	emergent adverse events in pediatric patier treated with Daytrana TM for 12 hours daily we anorexia (87 subjects, 46%), Insomnia (57 su jects, 30%), viral infection (54 subjects, 28%) awn from the study because of treatment-emerge
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UNUS AUSE AND UPPENDENCE Controlled Substance Class: DaytranaTM (methylphenidate transdermal system), like other methylphenidate products, is classified as a Schedule II controlled substance by federal regulation. Abuse, Dependence, and Tolerance: See WARNINGS-Drug Dependence for boxed warning containing drug abuse and dependence information. OVERDOSAGE

dependence information. **OVERDOSAGE** Signs and Symptoms: Signs and symptoms of acute methylphenidate overdosage, resulting principally from overstimulation of the CNS and from excessive sympathomimetic effects, may include the following: vomiting, aglatation, tremors, hyperreflexia, muscle twitching, convulsions (may be followed by corna), euphoria, contuison, hallucinations, delinum, sweating, flushing, headache, hyperpryrxia, tachycardia, palpitations, cardiac arrhythmias, hypertension, mydriasis, and dryness of muscle muscle twitching convulsions (may be followed by corna), euphoria, contusion, hallucinations, delinum, sweating, flushing, headache, hyperpryrxia, tachycardia, palpitations, cardiac arrhythmias, hypertension, mydriasis, and dryness of muscle twitching adhesive. The continuing absorption of methylphenidate from the skin, even after removal of the patch, should be considered when treating patients with overdose. Treatment consists of appropriate supportive measures. The patient must be protected against self-ficacy of periodenal dialysis or extracorporeal hemodalysis for Daytran⁴⁷ overdosage has not been estabilished. Poison Control Center, As with the management of all overdosages, the possibility of multiple drug ingestion should be con-didered. The physician may wish to consider contacting a poison control center for up-to-date information on the management of overdosage with metrylphenidate. Do not store patches unpouched. Store at 25 C (77 F); excursions permitted to 15-30° C (59-86° F) [see USP Controlled Room Temperature]. Unce the tray is opened, use contentis within 2 months. Apply the patch immediately upon removal from the protective pouch. Do not store patches unpouched. For transfermature only.

REFERENCE American Psychiatric Association. Diagnostic and Statistical Manual of Mental Disorders. 4th ed. Washington, DC: American Psychiatric Association 1994. Manufactured for Shire US Inc., Wayne, PA 19087 by Noven Pharmaceuticals, Inc., Miami, FL 33186. For more information call 1-400-282-2088 or visit <u>www.shire.com</u>. Dot Matrix[™] is a trademark of Noven Pharmaceuticals, Inc. Daytrana[™] is a trademark of Shire Pharmaceuticals, Inc. @ 2006 Shire Pharmaceuticals Ireland Limited.

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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.

Break the cycle

recurrence

residual symptoms sadness low energy anxiety

of unresolved depression with EFFEXOR XR",2

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, aggressiveness, impulsivity, akathisia, hypomania, and mania have been reported and may represent precursors to emerging suicidality. Stopping or modifying therapy should be considered especially when symptoms are severe, abrupt in onset, or not part of presenting symptoms.

relapse

- Treatment with venlafaxine is associated with sustained increases in blood pressure (BP) in some patients. Postmarketing cases of elevated BP requiring immediate treatment have been reported. Pre-existing hypertension should be controlled. 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.



Please see brief summary of Prescribing Information on adjacent pages.



BRIEF SUMMARY. See package insert for full prescribing information

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 psychlatric 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), obsessive-compulsive disorder (0CD), 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.

suicides occurred in these trials. CONTRAINDICATIONS: Hypersensitivity to veniafaxine hydrochloride or to any excipients in the formulation. Concomitant use in patients taking monoamine oxidase inhibitors (MADIS). WARNINGS: Clinical Worsening and Suicide Risk— Patients with major depressive disorder (MDD), both adult and pediatric, may experience worsening of their depression and/or the emergence of suicidall ideation and behavior (suicidality) or unusual changes in behavior, whether or not they are taking antidepressant medications, 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 depression and/or the emergence of suicidall vin certain patients. Antidepressants increased the risk of suicidal thinking and behavior (suicidality) in certain patients. Antidepressants increased the risk of suicidal thinking and behavior (suicidality) in short-term studies in children and adolescents with MDD and other psychiatric disorders. It is unknown whether the suicidality risk in pediatric patients extends to longor-term use, i.e., beyond several months. It is also unknown whether the suicidality risk extends to adults. All pediatric patients being threapy, or at times of dose changes, either increases or decreases. Adults with MDD or comorbid depression in the setting of other psychiatric linecaeses impulsivity, axathisia (psychomotor restlessness), hypomania, and mania have been reported in adult and norrasychiatric. Although a causal link between the emergence of such symptoms and either the suicidality represents with antidepressents should be observed similarly for clinical worsening and suicidality especially during the initial few months of a course of drug therapy, or at times of dose changes, either increases or decreases. Anxiety, agliation, panic attacks, insomnia, initability, hostily, aggressiveness impulsivity, axathisia (psychomotor restlessness), hypomania, and mania have been r Increases or decreases. Anxiety, agitation, panic attacks, insomila, intability, hostility, agressiveness, impulsivity, akathisa (goychomor resitessness), hypornania, and main have been reported in adult and pediatric patients being treated with antidegressants for MDD and other indications, both psychiatric and orpsychiatric. Although a causal in botween the emergence of suck wymittoms and efficient the worsening of depression and/or the emergence of sucked impulses has not been established, there is concern that such worsening depression or sucked in the decision has been made to discontinue treatment, medication persistently worse, or who are acperiencing emergent sucked any construct or sets or were not part or the patient's presenting symptoms. If the decision has been made to discontinue treatment, medication or notation generated as anyloid us its sabilis, but with recognition that attory discontinue treatment, medication of pediatric patients being treated with antidepressants for MDD or other indications, both psychiatric and nonpsychiatric, should be alerted about the need to monitor patients for the emergence of pediatric patients being treated with antidepressants for MDD or other indications, both psychiatric and nonpsychiatric, should be alerted about the need to monitor patients for the same sec-sticule the risk of overdose. Families and caregivers of adults being treated for depression should be similarly valveds. Careening Patients for Biolar Usoder: A major depression elevated the similar y advised. Screening Patients for Biolar Usoder: A major depression elevated be similarly valveds. Careening Patients for Biolar Usoder: A major depression elevated be aligned advised. Screening Patients for Biolar Usoder: A major depression is unknown. Prior to Initiation of aligned should be aligned baboer represent such a conversion is unknown. Prior to Initiation of aligned baboer and advise strengtons, and baboer secrets the depression is unknown. Prior to Initiation of aligned baboer

stadis. The decontrustion rate for arounds was 1/26 in MOD stadies. The decontrustion rate more formorely records for Effect AT (5%) then people (7%) plants in OD stadies. The decontrustion rate for the provide and Color than people controls was and the control in position of the control in the filter AT (Col) fan pieceb CPA (10%) that is the stade of the control in position of the control in the plants and the control in the plants in the stade of the control in position of the control in the plants and the control in the plants in the plants in the stade of the control in the plants and the control in the plants in the plants in the plants and the the control in the plants and the plants and the plants in the plants in the plants and the states and plants and the plants and the plants in the plants in the plants and the states and plants and the plants and the plants and the plants and the states and plants and the plants and the plants and the plants and the states and plants and the plants and the plants and the plants and the states and plants and the plants and th

studies. The discontinuation rate for anorexia was 1.0% in MDD studies. Treatment-emergent anorexia was more commonly reported for Effexor XR (8%) than placebo (2%) patients in GAD studies. The discontinuation rate for anorexia was 0.9% for up to 8 weeks in GAD studies. Treatment-emergent anorexia was more commonly reported for Effexor XR (20%) than placebo (2%) patients in SAD studies. The discontinuous nate for anorexia was 0.4% for up to

vasodilatation, thinking abnormal, decreased libido, and sweating. *Commonly Observed Adverse Events in Controlled Clinical Trials for MDD, GAD, SAD, and PD*—Body as a Whole: asthenia, headache, flu syndrome, accidental injury, abdominal pain. *Cardiovascular*: vasodilatation, hypertension, palpitation. <u>Digestive</u>: nausea, constipation, anorexia, vomiting, flatulence, diarrhea, eructation. *Metabolic/Nutritional:* weight loss. *Nervous System*: dizziness, somnolence, insomnia, dry mouth, nervousness, abnormal dreams, tremor, depression, hypertonia, paresthesia, ibido decreased, agitation, ankiety, twitching. <u>Respiratory System</u>: abnormal disculation, impotence, orgasmic dysfunction (including anorgasmia) in females. *Vital Sign Changes*: Effevor XR was associated with a mean increase in pulse rate of about 2 beats/min in depression and GAD trials and a mean increase in pulse rate of a beats/min in SAD trials. (See WARNINGS-Sustained Hypertension). *Laboratory Changes*: Clinically relevant increases in serum cholesterol were noted in Effevor XR clinical trials. Increases Changes: Clinically relevant increases in serum choesterol were noted in Finkon An clinical trads. Increases were duration dependent over the study period and tended to be greater with higher doses. **Other Events Observed During the Premarketing Evaluation of Effexor and Effexor XR**—N=6,670. "Frequent"=events occurring in at least 1/100 patients; "infrequent"=1/100 to 1/1000 patients; "rare"=fewer than 1/1000 patients. **Body as a whole** - Frequent: chest pain substernal, chills, fever, neck pain; Infrequent face edema, intentional injury, malaise, moniliasis, neck rigidity, pelvic pain, photosensitivity reaction, suicide attempt, withdrawal syndrome; Rare: appendicitis, bacteremia, carcinoma, cellulitis, **Cardiovascular system** - Frequent: <u>every of the sectural burgers of the secture of the sect</u> Intentional injury, malaise, moniliasis, neck rigidity, pelvic pain, photosensitivity reaction, suicide attempt, withdrawal syndrome; Rare: appendicitis, bacteremia, carcinoma, cellulitis. **Cardiovascular system** - Frequent: migraine, postural hypotension, tachycardia; Infrequent: angina pectoris, arrhythmia, extrasystoles, hypotension, peripheral vascular disorder (mainly cold feet and/or cold hands), syncope, thrombophilebitis; Rare: aortic aneurysm, arteritis, first-degree atrioventricular block, bigeminy, bundle branch block, capillary fragility, cerebral ischemia, coronary artery disease, congestive heart failure, heart arrest, hematoma, cardiovascular disorder (mirital valve and circulatory disturbance), muccottaneous hemorrhage, myocardial infarct, pallor, sinus arrhythmia. **Digestive system** - Frequent: increased appetite; infrequent: bruxism, colitis, rectal hemorrhage, hemorrhoids, melena, oral moniliasis, stomatitis, mouth ulceration; Rare: abdominal distension, biliary pain, cheilitis, cholecystitis, cholelithiasis, esophageal spasms, duodenitis, lietlis, jaundice, intestinal obstruction, liver tendemess, parotitis, periodontitis, protis, rectal disorder, salivary gland enlargement, increased salivation, soft stools, tongue discoloration. Endocrine system - Frae: galacorrhoea, golier, hyperthyroidism, hypothyroidism, thyroid nodule, thyroiditis. **Hemic and Lymphatic system** - Frequent: echymosis; Infreguent: amenia, leukocytosis, leukopenia, Lymphadenopathy, thrombocythemia, Rare: bhosphataes increased, diabylacia and **nutritional** - Frequent: edema, weight gain, Infreguent: alkaline phosphataes increased, dehydration, hyperchioesteremia, hyperglycemia, hyperfilemia, hypoglycemia, hyposhatemia, SGOT increased, SCPT increased, thirst; Rare: alcohol intolerance, bilirubinemia, hypothypotise, humer thesis, hypercholesteremia, hyperpenhosphatemia, hypothypotise, humer thesis, hypercholesteremia, hyperphosphatemia, hypothypotise, homer the solution, obschuld, unite file fasted. Application is performed weight manyer inference indexing and the product and the solution in the performant of the solution in the second solution is and in the second solution in the second solution is and solution in the second solution in the second solution is and solution in the second solutis in the se

Take a closer look at Dialogues Time to Talk

Dialogues

is a unique patient support and education program that is designed to help you foster successful therapy

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offers patients access to a call center to speak with a health care provider for patient support and education to reinforce your efforts

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The most common adverse events reported in EFFEXOR XR short-term placebo-controlled depression, generalized anxiety disorder (GAD), social anxiety disorder (SAD), and/or panic disorder (PD) 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.



The change they deserve.

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

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- Identification of Common Genetic Traits Across Different Disorders May Improve Treatment
- Paroxetine May Benefit Patients with Dysthmic Disorder
- History of Past Traumatic Incidents not Increased Among OCD Patients
- Schizophrenia May be More Severe in Patients with Tardive Dyskinesia

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Dan J. Stein, MD, PhD, University of Cape Town; Mark Solms, PhD, University of Cape Town; and Jack van Honk, PhD, Utrecht University

CME QUIZ

642 The quiz is CME-accredited by Mount Sinai School of Medicine for 3.0 credit hours.

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I never thought I could be myself again Now I can

Now the most prescribed atypical*

Proven efficacy To help patients achieve continued success¹¹⁻⁴

To help patients stay on treatment¹⁻⁵

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.

Elderly patients with dementia-related psychosis treated with atypical antipsychotic drugs are at an increased risk (1.6 to 1.7 times) of death compared to placebo (4.5% vs 2.6%, respectively). SEROQUEL is not approved for the treatment of patients with dementia-related psychosis.

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.

*All atypical prescriptions: Total prescriptions. Jan. 05-Feb. 06. New prescriptions. Sept. 04-Feb. 06. IMS Health. National Prescription Audit.

Significant improvement in all 11 YMRS items was measured at Day 21 and continued through Day 84 in monotherapy mania trials.

Please see Brief Summary of Prescribing Information on adjacent page.

5 Seroquel quetiapine fumarate

25 mg, 50 mg, 100 mg, 200 mg, 300 mg & 400 mg tablets

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es: 1. Vieta E, Mullen J, Brecher M, et al. Quetiapine monotherapy for mania associated with bipolar disorder: combined analysis of two international, double-blind, randomised, placebo-controlled studies. *Curr Med Res Opin*. 2005;21:923-934. G, Chengappa KNR, Suppes T, et al. Quetiapine with libium or divalpreex for the treatment of bipolar mania: a randomized, double-blind, placebo-controlled study. *Bipolar Disord*. 2004;6:213-223. **3**. Small JG, Kolar MC, Kellams JJ. et in achizophrenia: onset of achizophremia: onset of achizophremia. *Curr Med Res Opin*. 2004;20:107-1023. **4**. Kasper S, Brecher M, Fitton L, et al. Maintenance of long-term efficacy and safety of quetiapine in the open-label treatment of enia. *Int Clin Psychopharmacol*. 2004;19:281-289. **5**. SEROQUEL Prescribing Information.

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Processing momentum. Increased Mortality in Elderly Pallanta with Demantia-Relative Psychosis Ederly plation. Which doministra-ministed psychosis treatile with stypical antipsychotic drugs are at an increased rate of death compared to placeble, handpeer of severatem placebo-cantitulity of the cartifie of 10 weaks in these platents revealed after of calls in the drug-maid patients of between 1.8 to 1.7 times that seen in placeble-treated platinis. Duer the source of a typical 10 weak controlled weaks of the drug of the causes of death weaks of Artic 45%, compared to a not of about 2.5% in the placebo group. Although the causes of death weak version of the deaths appeared to be other car-dioversactive (synchratikines, under death) of indeficient (or go, poesumate) is notern. SEROUREL (quel-apine) is not approved in the treatment of pelients with Demantia-Related Psychosis.

Apriley is not approved for the treatment or petimena wink Leansance Treasmer + re-MINCATIONS AND USAGE: Deplar Mank: SERIOUEL is indicated for the treatment of acute marks episodes asso-ciated with block II disords, as ethnicities the monotherapy undirect threaty to Minlamo disordines. The effacts of SERIOUEL is name to block marks established in two 12-week monotherapy trials and one 5 -week adjunt threaty primal block and block marks and block marks and the series of the series

Contribution Carbon School Car

control to all relation on the delation of the strength generally, explored to particular to extend be detected and the strength of the str

sis, should lead to consideration of a lower starting does, glower titration, and careful monitoring during the initial does inge proof in the defay. The mang loarne of SEROULEL was reduced by 30% to 50% in defay pretents when compared to younger patients. AUVERSE REACTINGS: The information below is derived from a clinical trial detabase for SEROULEL consisting of over 3000 patients. This database includes 405 patients exposed to SEROULEL for the treatment of acute blogina main (anomatery and adjunct threapy) and approximately 2500 patients acyosed to 15 more doess of SEROULEL consisting of over 3000 patients. This database includes 405 patients exposed to 15 ROULEL for the treatment of acute blogina main (anomatery and adjunct threapy and approximately 2500 patients acyosed in 15 more doess of SEROULEL to the treatment of acute blogina provide the start of the treatment of schizophrenia. (Of these approximately 3000 aubjects, approximately 200 (2001 ns shortphrenia and 405 match bagina main). Adverse Flatingto Seroever in Sikof-Team, Platebo-Controlled Triats: Reptra Mantic - Outeral, discontinuations of treatment in Sikof-Team, Platebo-Controlled Triats: Biplard Mantic - Outeral, discontinuations due to source events verte 5, % for SEROULEL s. 51% to placebo in montherapy and 30% for SEROULEL s. 55% or placebo and input therapy. Schloghennia: Overal, there was also discontinuation due to adverse events (% for S SeROULEL s. 51% to placebo in apoid for controlled Triats. The prescription e0.0% or 5% or placebo and input the sites and tabulations in the full apoid to controlled Triats. The second source e0.5% or 5% replacebo and input the sites and the sites and tabulations and approximation cannot be used to predict the index deve the disk in the Sites of clausions in the full Prescription Information cannot be used to predict the index of site of thesis in the course of usual mulcial prescription information cannot be used to predict the index of the index of the bloging information disk of t

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