CNS SPECTRUMS® The International Journal of Neuropsychiatric Medicine



Index Medicus/MEDLINE citation: CNS Spectr



NEW ADULT INDICATION

Because he's in

Aim Higher With ADDERALL XR[®] —

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

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

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

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

www.ADDERALLXR.com www.ADHDSupportCompany.com



demand all day long...

AT WORK For Efficacy That Measures Up to Life's Demands

- Once-daily dosing provides all-day symptom control'
- Mean ADHD-RS total scores for adults receiving **ADDERALL XR** decreased by 41%¹
- ADDERALL XR is the only stimulant medication approved to treat adults with ADHD¹
- Clinical data in adults demonstrate that **ADDERALL XR** is generally well tolerated¹





References: 1. Ambrosini PJ, Lopez FA, Chandler MC, et al. Safety and efficacy of ADDERALL XR in pediatric ADHD: results of an open-label community assessment trial. Poster presented at: 14th Annual CHADD International Conference October 17, 2002; Miami Beach, Fia. 2. Spencer T, Biederman J, Wiens T, et al. Pharmacotherapy of attention-deficit. hyperactivity disorder across the life cycle. J Am Acad Child Adoles: Psychiatry, 1996;35:409-432. 31. 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 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; Lopez FA. Chandler MC, Biederman J, et al. Long-term Adderall XR treatment improves quality of life in ADHD ADHD. International Meeting of the American Psychiatric Association, May 21, 2003; San Francisco, Calif. 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

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

Advanced arteriosclerosis, symptomatic cardiovascular disease, moderate to severe hypertension, hyperthyroidism, known hypersensitivity or idiosyncrasy to the sympa-tomimetic 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 ic use of stimulants in children, including amphetamine, may be causally associated with suppression of growth. Therefore, growth should be monitored during treatment, and patients who are not growing or gaining weight as expected should have their treatment interrupted. Sudden Death and Pre-existing Structural Cardiac Abnormalities: Sudden death has been reported in association with amphetamine treatment at usual doses in children with structural cardiac abnormalities. Adderall XR® generally should not be used in children or aduits with structural cardiac abnormalities.

PRECAUTIONS

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

General: The least amount of amphetamine feasible should be prescribed or dispensed at one time in order to minimize the possibility of overdosage. Hypertension: Caution is to be exercised in prescribing amphetamines for patients with even mild hypertension (see CONTRANUCIATIONS). Bilood pressure and pulse should be monitored at appropriate intervals in patients taking ADDERALL XRP, especially patients with hypertension. Ties: Amphetamines have been reported to exacerbate motor and phonic tics and Tourette's syndrome. Therefore, clinical evaluation for tics and Tourette's syndrome in children and their families should precede use of stimulant medications.

Ites: Amphetamines have been reported to exacerbate motor and phonic tics and Tourette's syndrome. Therefore, clinical evaluation for tics and Tourette's syndrome in children and their families should precede use of stimulant medications. Information for Patients: Amphetamines may impair the ability of the patient to engage in potentially hazardous activ-rities such as operating machinery or vehicles; the patient should therefore be cautioned accordingly. Drug Inferentions: Activitying agents—Gastrointestinal activitying agents—These agents (ammoni-um choride, sodium acid phosphate, etc.) increase the concentration of the ionized species of the amphetamine molecule, thereby increasing unitary excretion. Both groups of agents lower blood levels and efficacy of ampheta-amines. Adrenergic blockers—Adrenergic blockers are inhibited by amphetamines. Adrenergic blockers—Adrenergic blockers are inhibited by amphetamines. Adrenergic blockers—Adrenergic blockers are inhibited by amphetamines. Advanta akainizing agents (sactazotamide, some thaizdes) increase the concentration of the non-ionized species of the amphetamine molecule, thereby decreasing urnary excretion. Both groups of agents increase blood levels and therefore potentiate the actions of amphetamines. Anticepressants, truc/clio—Amphetamines may enhance the activity of tricyclic antidepressants or sympathominetic agents; d-amphetamines, increase blood levels and therator potentiate the activity of tricyclic antiphetamines, antietasing their effect on the relaxe of norepinephrine and other monoamines from adrenergic increases in the concentration of d-amphetamine in the brain; cardiovascular effects can be potentiated. *MAO inhibitors*—MAOI antidepressants, as well as a metabolite on the relaxe of norepinephrine and other monoamines from adrenergic nerve endings; this can cause beadches and other signs of hypertensive crisis. A variety of toxic neurological effects and maignant hyperprytexia can courdovascular effects danabetamines may be inhibi

mine CNS stimulation is potentiated and fatal convulsions can occur. Veratrum alkaloids—Amphetamines inhibit the hypotensive effect of veratrum alkaloids. Drug/Laboratory 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 Ferlilly: No evidence of carcinogeneity was found in studies in which d.l-amphetamine (enantiomer ratio of 1:1) was administered to mice and rats in the diet for 2 years at doses of up to 30 mg/kg/day in male mice, 19 mg/kg/day in female 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/day [child] on a mg/m' body surface area basis. Amphetamine, in the enantiomer ratio present in ADDERALL® (immediate-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 *E. coli* component of the Ame stet *in vitro*. 0.4-Amphetamine (1:1 enantiomer ratio) has been reported to produce a positive response in the *invitro* Sister chromatid exchange and chromosomal aberration assays. Amphetamine, in the enantiomer ratio present in ADDERALL® (immediate-release) (d- to 1- ratio of 3:1), did not adversely affect fertility or early embryonic development in the rat at doses of up to 20 mg/kg/day (approximately 5 times the maximum recommended human dose of 30 mg/day on a mg/m' body surface area basis). **Pregnancy:** Pregnancy Category C. Amphetamine, in the enantiomer ratio present in ADDERALL® (de 1- ratio of 1:1), had not adversely affect fertility or early embryonic development or survival when orally administered to pregnant rats and rabbits throughout the period of organogenesis at doses of up to 6 and 16 mg/kg/day (approximately 5 times that of a human dose of 30 mg/day (ohild) on a mg/m' body surface area b

A number of studies in rodents indicate that prenatal or early postnatal exposure to amphetamine (d- or d,I-), at doses similar to those used clinically, can result in long-term neurochemical and behavioral alterations. Reported behavioral affects include learning and memory deficits, affered locomotor activity, and changes in sexual function. There are no adequate and well-controlled studies in pregnant women. There has been one report of severe congenital bony deformity, trackeo-esophageal fistula, and anal atresia (vater association) in a baby born to a woman who took dextroamphetamine sulfate with lovastatin during the first trimester of pregnancy. Amphetamines should be used during pregnancy only if the potential benefit justifies the potential its to the fetus. Nonteratogenic Effects: Inducts born to mothers dependent on amphetamines have an increased risk of premature delivery and low birth weight. Also, these infants may experience symptoms of withdrawal as demonstrated by dysphoria, including apitiation, and significant lassitude. **Usage In Nursing Mothers:** Amphetamines are excreted in human milk. Mothers taking amphetamines should be advised to refrain from nursing. **Perilatic Use:** ADDERALL XR® is indicated for use in children 6 years of age and older. **Usa in Children Under Six Years of Age:** Effects of ADDERALL XR® in 3-5 year olds have not been studied. Long-term effects of amphetamines are to thildren have not been well established. Amphetamines are not recommended for use in children under 3 years of age. **Geriatric Use:** ADDERALL XR® has not been studied in the geriatric population. **ADVERSE EVENTS**

ADVERSE 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

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5 mg, 10 mg, 15 mg, 20 mg, 25 mg, 30 mg CAPSULES (Mixed Salts of a Single-Entity Arnphetamine Product) Dextroamphetamine Sulfate Dextroamphetamine Sacharate Amphetamine Aspartate Monohydrate Amphetamine Sulfate

Site presented at: 15th Annual Meeting or the American registration association, my an average and the association of the anti-case of the association of the associa standardized events a legunes. In the tables and instings that follow, cost and reliminology has been used to classify reported adverse events. The stated frequencies of adverse events represent the proportion of individuals who experienced, at least once, a treatment-emergent adverse event of the type listed.

Adverse events associated with discontinuation of treatment: In two placebo-controlled studies of up to 5 weeks duration among children with ADHD, 2.4% (10/425) of ADDERALL XR® treated patients discontinued due to adverse events (including 3 patients with loss of appetite, one of whom also reported insomnia) compared to 2.7% (7/259) receiving placebo. The most frequent adverse events associated with discontinuation of ADDERALL XR® (ontrolled and uncontrolled, multiple-dose clinical triats of pediatric patients (Ne595) are presented below. Over half of these patients were exposed to ADDERALL XR® for 12 months or more.

% of pediatric patients discontinuing (n=595) 2.9 1.5 1.2

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

0.7 In one placebo-controlled 4-week study among adults with ADHD, patients who discontinued treatment due to adverse events among ADDERALL XR®-treated patients (N=91) were 3.1% (n=6) for nervousness including anxiety and irritability. 2.6% (n=5) for insomnia, 1% (n=2) each for hat increase, apitation, chest pain, cocaine craving, elevated blood pressure, and weight loss.

10

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 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 course of usual medical practice where patient characteristics and other factors differ from those which prevailed in the clinical trials. Similarity, 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.

Body System	Preferred Term	ADDERALL XR® (n=374)	Placebo (n=210)
General	Abdominal Pain (stomachache)	14%	10%
	Accidental Injury	3%	2%
	Asthenia (fatigue)	2%	0%
	Fever	5%	2%
	Infection	4%	2%
	Viral Infection	2%	0%
Digestive	Loss of Appetite	22%	2%
System	Diarrhea	2%	1%
	Dyspepsia	2%	1%
	Nausea	5%	3%
	Vomiting	7%	_4%
Nervous System	Dizziness	2%	0%
	Emotional Liability	9%	2%
	Insomnia	17%	2%
	Nervousness	6%	2%
Metabolic/Nutritional	Weight Loss	4%	0%

Table 2 Adverse Events Reported by 5% or More of Adults Receiving ADDERALL XR® with Higher Incidence

Body System	Preferred Term	ADDERALL XR® (n=191)	Placebo (n=64)
General	Asthenia Headache	6% 26%	5% 13%
Digestive System	Loss of Appetite Diarrhea Dry Mouth Nausea	33% 6% 35% 8%	3% 0% 5% 3%
Nervous System	Agitation Anxiety Dizziness Insomnia	8% 8% 7% 27%	5% 5% 0% 13%
Cardiovascular System	Tachycardia	6%	3%
Metabolic/Nutritional	Weight Loss	11%	0%
Urogenital System	Urinary Tract 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, photosensilivity reaction, constpation, note in disorder, emotionai lability, libido decreased, somnolence, speech disorder, patipation,

reaction, constipation, tooth disorder, emotional lability, libido decreased, somnolence, speech disorder, palpitation, wriching, dyspina, sweating, dysmenorrhea, and impotence. "included doses up to 60 mo, 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, dystemesia, dysphoria, depression, tremor, headache, exacerbation of motor and phonic tics and Tourette's syndrome, seizures, struck Gastrointestinal: Dryness of the mouth, unpleasant taste, diarrhea, constipation, other gastrointestinal disturbances. Anorexia and weight loss may occur as undesirable effects. Allergic: Unicaria. Endocrine: Impotence, changes in libido. **DRUG ABUSE AND DEPENDENCE** ADDERALL XMP is a Schedule II controlled substance.

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.

OVERDOSAGE

from schizophrena. **OVERDOSAGE** Individual patient response to amphetamines varies widely. Toxic symptoms may occur idiosyncratically at low doses. Symptoms: Manifestations of acute overdosage with amphetamines include restlessness, tremor, hyper-reflexia, rapid respiration, confusion, assaultiveness, hallucinations, panic states, hyperpyrexia and rhabdomyol-ysis. Fatigue and depression usually follow the central nervous system stimulation. Cardiovascular effects include arrhythmias, hyperension or hypotension and circulatory collapse. Castrointestinal symptoms include nausea, vomiting, diarrhea, and abdominal cramps. Fatal poisoning is usually preceded by convulsions and come amphetamine intoxication is largely symptomatic and includes gastric lavage, administration of activated charcoal, administration of a cathartic and sedation. Experience with hemodialysis or peritoneal daiysis is inadequate to increase risk of acute renal failure if myoglobinuria is present. If acute severe hypertension complicates amphetamine stimulant effects of amphetamines and can be used to treat amphetamine intoxication. The prolonger release of may and the submit control acute at the severe hypertension complicates ampheta-mine overdosage, administration of intravenous phentolamine has been aclieved. Chlorpromazine antagonizes the central stimulant effects of amphetamines and can be used to treat amphetamine intoxication. The prolonger release of mixed amphetamine safts from ADDEFALL XR® should be considered when treating patients with overdose. Dispense in a tight, light-resistant container as defined in the USP. Store at 25° C (77° F). Excursions permitted to 15° So² C (59-6F f) [see USP controlled Room Temperature]. Manufactured for: **Shire US Inc.**, Newport, KY 41071 Made in USA For more information call 1-800-828-2088, or visit www.adderalkr.com. ADDEFALL² and ADDEFALL XR® are registered in the USP Store at 25° C (77° F). Excursions permitted to 15° Sire US Inc. Newport, KY 41071 Made in USA For more

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Rev. 9/04





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Available online at www.adaa.org

CAREER DEVELOPMENT TRAVEL AWARD - \$3,500 Deadline: Tuesday, November 30, 2004

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

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

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

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

About the ADAA

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

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

BRIEF SUMMARY of PRESCRIBING INFORMATION INDICATIONS AND USAGE: Bipolar Mania: SEROQUEL is indicated for the short-term treatment of acute manic INDICATIONS AND USACE: Bigdar Maria: SEROUEL is indicated for the short-term resument of acute manic epostos associaties with bejord indicatorie as ether monotherapy or diginut hetapy to fibitum of divergors. The efficacy of SEROUEL in acute biopiar mania was established in you Sweek monotherapy traits and one 3-week adjunct therapy truth of bopiar i plateini trildally hospitaled for up to 7 days for caute manic. Efficiencess for more than 3 weeks has not been systematically exultated in clinicial trials. Therefore, the physician who elects to us SEROUEL in ordered periods should periodically re-evaluate the long-term rises and benefit or the drug for the individual plateint. Sehizophrenia: SEROUEL is indicated for the trastment of schizophrenia. The efficiary of SEROUEL in conclusion in the senior of SEROUEL in order the senior of schizophrenia. The efficiary of evaluated in controller trials of schizophrenia weeks has not been systematically evaluated in controller to its schizophrenia was that is, for more than 6 weeks, nos not been systematically evaluated in controller to its business the physican who devices to use SEROUEL is ordered periods should periodically re-evaluate the long-term usefulness of the drug for the individual patient.

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

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WARNERS: Neuropean Selfcers Magnet Synchrone (MSS) is better reported to RSPCULE. Their and the implementation of artisphoto grant data selfcers in a section with antimetation of artisphoto grant data selfcers. The implementation of artisphoto grant data selfcers and annual selfcers and annual selfcers and annual selfcers. The implementation of artisphoto grant data selfcers and annual selfcers and annual selfcers and annual selfcers. The implementation of artisphoto grant data selfcers and annual selfcers and annual selfcers and annual selfcers. The implementation of annual selfcers and annual selfcers annual selfcers and annual selfcers and annual selfcers and annual selfcers annual serum protach i week is unknown for most paliests. Neither chical is dudes nore potentially be explored in the conductive at this time. Transmisses Elevations of the conductive at this time transmisses elevations of the conductive at this time. Transmisses Elevations of the conductive at the conductive at this time transmisses elevations of the conductive at the con

SEROQUEL® (quetiapine furnarate) Tablets

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Steven Johnson syndrome (SUS). **DRUG ABUSE AND DEPENDENCE:** Controlled Substance Class: SERGOUEL is not a controlled substance. Physical **and Psychologi Exemptions:** SERGOUEL has not been systematically slutied, in minats or humans, for its politi-fail for allow, tolerance or physical dependence. When the clinical traits during the original dependence and the systematically is not possible to prevent any tandency for any droug-besing behavior, these sobservations were the systematically the physical or the table of the limit of experiments the estent to which a XIS-active drop will be missaed, develop and we show there manded Depared to physical dependence cardinuly for a hybrid to and they during and such attaines should be observed to develop for signs of missae or abuse of SERGOUEL, e.g., development of tolerance, microases in doze, drog-seeding behavior.

OVERDOSAGE: Human experience: Experience with SEROQUEL (quetiapine furnarate) in acute overdosag verouosad: numan apprenting. Experience with Schoolder (participite fundate) in Abite Verousage was limited in the clinical trial database (6 reports) with estimated doses ranging from 1200 mg to 9600 mg and no fabilities in general, reported signs and symptoms were those resulting from an exaggregation of the drug known pharmacological effects, i.e., drowsness and sedation, tachycardia and hypotension. One case, involving In leases in general, reported spice and symptoms were most resumption an escaperation of the study sown pharmacobipatie effects, i.e., downess and seation, halkhyada and hybotesion. One case, involving an estimated overdose of SROV may associated with hybotelemic and first degive herat block in post-neting caperianc. The harbe ener very structures of SROVDEL and the resulting in a share the off a source degiuse to operation and vertices age. The source of seating caperiance in the structure of seating caperiance. The source of seating caperiance is the source of seating caperiance in the source seating in a structure of the source of seating caperiance. The source of seating caperiance is the source of seating caperiance is and vertices of a source degiuse is a result of a source degiuse caperiance match the and in nokel distortion (we result in a structure of seating caperiance) is and vertices of seating caperiance in the source of seating caperiance is and vertices of seating caperiance. The possibility of obtitude included the seating caperian results of the source of seating caperian caperian seating and the seating caperian caperian seating caperian in the source of seating caperian caperians and the source of seating caperians of the seating caperian caperians in the source of seating caperians and vertices and caperians the source of seating caperians and caperians that caperians the source of the seating caperians and caperians that caperians that caperians that caperians that caperians the source of the seating caperians and caperians that caperians that caperians the source of the seating caperians and caperians that the structure of the structure of the s SERODUEL is a trademark of the AstraZeneca group of companies.

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Rev 01/04

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I never thought I could be myself again

Now I can

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

Eno

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

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

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



AstraZeneca Pharmaceuticals LF

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Table of Contents

- 803 Introduction: The Multiple Facets of Treatment-Resistant Depression Daniel Souery, MD, PhD, Erasme Hospital; and Karel Van der Auwera, Erasme Hospital
- Therapeutic Alternatives for Difficult-to-Treat Depression: 808 A Narrative Review of the State of the Evidence Michael E. Thase, MD, University of Pittsburgh Medical Center
- 823 Therapy of Treatment-Resistant Depression: Focus on the Management of TRD with Atypical Antipsychotics Nikolas Klein, MD, Medical University of Vienna; Julia Sacher, MD, Medical University of Vienna; Helene Wallner, MD, Medical University of Vienna; Johannes Tauscher, MD, Medical University of Vienna; and Siegfried Kasper, MD, Medical University of Vienna
- 833 A Review of the Treatment for Refractory Obsessive-Compulsive Disorder: From Medicine to Deep Brain Stimulation David S. Husted, MD, University of Florida College of Medicine; and Nathan A. Shapira, MD, PhD, University of Florida College of Medicine
- 849 The Role of Atypical Antipsychotics in Glucose/Insulin Dysregulation and the Evolving Role of the Psychiatrist in a New Era of Drug Treatment Options Anthony Ferraioli, MD, Shaker Park Medical; Kara Lee Shirley, PharmD, RPh, Albany College of Pharmacy; and Panakkal David, MD, Albany Medical College
- 862 Clinical Experience with Aripiprazole Treatment in Ten Elderly Patients with Schizophrenia or Schizoaffective Disorder: Retrospective Case Studies Subramoniam Madhusoodanan, MD, St. John's Episcopal Hospital; Ronald Brenner, MD, St. John's Episcopal Hospital; Sanjay Gupta, MD, Olean General Hospital; Harsha Reddy, MD, St. John's Episcopal Hospital; and Olivera Bogunovic, MD, T.L.C. Health Care Network

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.

CNS SPECTRUMS The International Journal of Neuropsychiatric Medicine

Table of Contents

Departments/Monthly Columns

FROM THE EDITOR'S DESK

796 Ever-Present Adverse Effects By Jack M. Gorman, MD

CLINICAL UPDATES IN NEUROPSYCHIATRY

797 Breaking News From the Field of Neuroscience

- FDA Issues Black Box Warning for All Antidepressants
- Clozapine and Olanzapine May Increase Risk of Diabetes for Patients with Schizophrenia
- Parkinson's Disease and Epilepsy Linked to Depression

News From the 17th Congress of the European College of Neuropsychopharmacology

- Adjunctive Modafinil May Improve Fatigue and Depressive Symptoms in Major Depressive Disorder
- Quetiapine May Be Effective in Treating Bipolar I Disorder
- Spanish Cross-sectional Study Examines Prevalence of Comorbidities in Patients with Schizophrenia

CME QUIZ

871 The quiz on treatment-resistant depression is CME-accredited by Mount Sinai School of Medicine for 3.0 credit hours.

ABRIDGED ACADEMIC SUPPLEMENT

Unmasking Bipolar Disorder: Overcoming the Barriers to Treatment Success By Roger S. McIntyre, MD, Jakub Z. Konarski, MSc, Prakash S. Masand, MD, Farhan S. Fazal, MBBS, Ashwin A. Patkar, MD, Michael W. Otto, PhD, and David J. Miklowitz, PhD Founded in 1996, *CNS Spectrums* is an *Index Medicus* journal and is available on MEDLINE under the citation *CNS Spectr.* It is available online at www.cnsspectrums.com.

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Abilify. Now Indicated For Bipolar Mania.

Imagine where this could lead

Abilify is indicated for the treatment of schizophrenia and acute manic and mixed episodes associated with bipolar disorder.

IMPORTANT SAFETY INFORMATION

As with all antipsychotic medications, a rare condition referred to as neuroleptic malignant syndrome (NMS) has been reported. As with all antipsychotic medications, prescribing should be consistent with the need to minimize the risk of tardive dyskinesia (TD).

Hyperglycemia, including some serious cases ranging from ketoacidosis to death, has been reported in patients treated with atypical antipsychotics. Abilify was not included in epidemiologic studies suggesting this risk; therefore the risk of hyperglycemia with Abilify is not known. However, there have been few reports of hyperglycemia in patients treated with Abilify. Patients should be appropriately tested before and monitored during treatment.

Abilify may be associated with orthostatic hypotension and should be used with caution in patients with known cardiovascular disease, cerebrovascular disease, or conditions which would predispose them to hypotension.

As with other antipsychotic drugs, Abilify should be used with caution in patients with a history of seizures or with conditions that lower the seizure threshold. Seizures occurred in 0.3% of bipolar patients treated with Abilify in placebo-controlled trials.

Bristol-Myers Squibb Company Otsuka America Pharmaceutical, Inc.

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©2004 Otsuka America Pharmaceutical, Inc., Rockville, MD D6-K0072 AP4537/09-04 October 2004 Patients should not drive or operate heavy machinery until they are certain Abilify does not affect them adversely.

Commonly observed adverse events reported with Abilify in 3-week bipolar mania trials at a \geq 5% incidence for Abilify and at a rate at least twice the rate of placebo include, respectively, akathisia (15% vs 4%), constipation (13% vs 6%), and accidental injury (6% vs 3%).

Treatment-emergent adverse events reported with Abilify in short-term trials at an incidence $\geq 10\%$ and greater than placebo, respectively, include headache (31% vs 26%), agitation (25% vs 24%), anxiety (20% vs 17%), insomnia (20% vs 15%), nausea (16% vs 12%), dyspepsia (15% vs 13%), somnolence (12% vs 8%), akathisia (12% vs 5%), lightheadedness (11% vs 8%), vomiting (11% vs 6%), and constipation (11% vs 7%).

The adverse events reported in a 26-week, double-blind schizophrenia trial comparing Abilify and placebo were generally consistent with those reported in the short-term, placebo-controlled schizophrenia trials, except for a higher incidence of tremor: 9% for Abilify vs 1% for placebo. In this study the majority of the cases of tremor were of mild intensity (9/13 mild and 4/13 moderate), occurred early in therapy (9/13 ≤49 days), and were of limited duration (9/13 ≤10 days). Tremor infrequently led to discontinuation

(<1%) of Abilify. In addition, in a long-term (52-week), active-controlled study, the incidence of tremor for Abilify was 4%.

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Brief Summary of Prescribing Information. For complete prescribing information please consult official package circular INDICATIONS AND LISAGE

Schizophrenia ABILIFY is indicated for the treatment of schizophrenia. The efficacy of ABILIFY in the treatment of schizophrenia was established in short-term (4- and 6-week) controlled trials of schizophrenic inpatients (see CLINICAL PHARMACOLOGY: Clinical Studies in Full Prescribing Information). The efficacy of ABILIFY in maintaining stability in patients with schizo-clinical Studies in Full prescribing Information). The efficacy of ABILIFY in maintaining stability in patients with schizo-term of the fact have prescribing information). The efficacy of ABILIFY in maintaining stability in patients with schizo-term of the fact have prescribed by stable on other antinsvchotic medications for periods of 3 months or longer, were Cuincial studies in full Preschong information). The emitcacy of ABuLIPY in maintaining stability in pagents with schizo-phrenia who had been symptomatically stabile on other antipsychotic medications for periods of 3 months or longer, were discontinued from those other medications, and were then administered ABILIPY 15 mg/day and observed for relapse dur-ing a period of up to 26 weeks was demonstrated in a placebo-controlled trial (see CLINICAL PHARMACOLOGY: Clinical Studies). The physician who elects to use ABILIPY for stended periods should periodically re-evaluate the long-term use-funces of the drug for the individual patient (see DOSAGE AND ADMINISTRATION in Full Prescribing Information). Bipolar Mania

Bipoter Mania ABILIPF is indicated for the treatment of acute manic and mixed episodes associated with Bipotar Disorder. The efficacy of ABILIPF was established in two placebo-controlled trials (3-week) of inpatients with DSM-IV criteria for Bipotar I Disorder who were experiencing an acute manic or mixed episode with or without psycholic features (see CLINICAL PHARMACOLOGY), however, the effectiveness of ABILIPF to tonger-term use, that is, for more than 3 weeks of treat-ment of an acute episode, and for prophytactic use in mania, has not been established in controlled clinical trials. Therefore, physicians who elect to use ABILIPF for extended periods should periodically re-evaluate the long-term risks and benefits of the drug for the individual patient (see DOSAGE AND ADMINISTRATION).

CONTRAINDICATIONS

ABILIFY is contraindicated in patients with a known hypersensitivity to the product.

WARNINGS

Meuroleptic Malignant Syndrome (NMS) A potentially fatal symptom complex sometimes referred to as Neuroleptic Malignant Syndrome (NMS) has been reported in association with administration of antipsychotic drugs, including aripiprazole. Two possible cases of NMS occurred dur-ing antiprazole treatment, in the premarketing worldwide clinical database. Clinical manifestations of NMS are hyperpyrexia, muscle rigidity, altered mental status, and evidence of autonomic instability (irregular pulse or blood pressure, tachycardia, diaphoresis, and cardiac dysritythmia). Additional signs may include elevated creatine phosphokinase, myo-giobinuri (hadodmoybis), and acute renal faiure. The diagnostic evaluation of patients with this syndrome is compligootime (macoorrigoysis), and actue retrainance. The badjuosic evaluation of patients with this synthetic is compli-cated. In arriving at a diagnosis, it is important to exclude cases where the clinical presentation includes both serous med-ical illness (e.g., pneumonia, systemic infection, etc.) and untreated or inadequately treated extrapyramidal signs and symptoms (EPS). Other important considerations in the differential diagnosis include central anticioningic toxicit) immediate stoke, drug fever, and primary coentral nervous system pathology. The management of NMS should include: 1) immediate discontinuation of antipsychotic drugs and other drugs not essential to concurrent therapy; 2) intensive symptomatic treatand a more than the second sec Tardive Dyskines

A syndrome of potentially irreversible, involuntary, dyskinetic movements may develop in patients treated with antipsy-chotic drugs. Atthough the prevalence of the syndrome appears to be highest among the elderly, especially elderly women, it is impossible to rely upon prevalence estimates to predict, at the inception of antipsychotic treatment, which values, it is impossible to by door heverate commarks to product, at the incipation of anticosyload to transfer, which patients are likely to develop the syndrome. Whether antipsychotic drug products differ in their potential to cause tardive dyskinesia is unknown. The risk of developing tardive dyskinesia and the likelihood that it will become inversible are believed to increase as the duration of treatment and the total cumulative dose of antipsychotic drugs administered to believed to increase as the duration of treatment and the total cumulative dose of antipsychotic drugs administered to the patient increase. However, the syndrome can develop, although much less commoniy, after relatively brief treatment periods at two doses. There is no known treatment for established cases of tardive dyskinesia, although the syndrome may remit, partially or completely, if antipsychotic treatment is withdrawn. Antipsychotic treatment, itself, however, may suppress (or partially suppress) the signs and symptoms of the syndrome and, thereby, may possibly mask the underly-ing process. The effect that symptomatic suppression has upon the long-term course of the syndrome is unknown. Given these considerations, ABILIFY should be prescribed in a manner that is most likely to minimize the occurrence of tardive dyskinesia. Chronic antipsychotic treatment should generally be reserved for patients who suffer from a chronic illness harmful treatments are not available or appropriate. In patients who do require chronic treatment, the smallest dose and the shortest duration of treatment producing a satisfactory clinical response should be acought. The need for continued treatment should be reassessed periodically. If signs and symptoms of tardive dyskinesia appear in a patient on ABILIFY, drug discontinuation should be considered. However, some patients may require treatment with ABILIFY despite the pres-ence of the syndrome. ence of the syndrome.

Perce or the syndrome. Hyperglycemia and Diabetes Mellitus Hyperglycemia, in some cases extreme and associated with ketoacidosis or hyperosmolar coma or death, has been reported in patients treated with atypical antipsychotics. There have been few reports of hyperglycemia in patients treated with ABILIPY, Although fewer patients have been treated with ABILIPY, it is nore limited experience is the sole reason for the paucity of such reports. Assessment of the relationship between atypical antipsychotic use and glucose sole reason for the paucity of such reports. Assessment of the relationship between atypical antipsychotic use and glucose abnormalities is complicated by the possibility of an increased background risk of diabetes melitius in patients with schiz-ophrenia and the increasing incidence of diabetes melitius in the general population. Given these confounders, the rela-tionship between atypical antipsychotic use and hyperglycemia-related adverse events is not completely understood. However, epidemiological studies which did not include ABLIPY suggest an increased risk of treatment-emergent hyperglycemia-related adverse events in patients treated with the atypical antipsychotics included in these studies. Because ABLIPY was not marketed at the time these studies were performed, it is not known if ABLIPY suggesthet with atypical antipsychotics are not available. Patients with an established diagnosis of diabetes mellitus who are started on atypical antipsychotics are not available. Patients with an established diagnosis of diabetes mellitus who are started on atypical antipsychotics are not available. Patients with an established diagnosis of diabetes mellitus who are started on atypical antipsychotics. antipsychotics are not available. Fatemits with an established braginoss of blacebes mentius who are stated of adplica-antipsychotics should be monitored regularly for worsening of glucose control. Patients with insis factors for diabetes mel-lifus (e.g., obesity, family history of diabetes) who are starting treatment with atypical antipsychotics should undergo fast-ing blood glucose testing at the beginning of treatment and periodically during treatment. Any patient treated with atypical antipsychotics should be monitored for symptoms of hyperglycemia including polydipsia, polyuria, polyuria, and weak-ness. Patients who develop symptoms of hyperglycemia during treatment with atypical antipsychotics should undergo fasting blood glucose testing. In some cases, hyperglycernia has resolved when the atypical antipsychotic was discontinued; however, some patients required continuation of anti-diabetic treatment despite discontinuation of the suspect drug.

PRECAUTIONS General

Orthostatic Hypotension

Orthostatic Hypotension Aripiprazole may be associated with orthostatic hypotension, perhaps due to its α_1 -adrenergic receptor antagonism. The incidence of orthostatic hypotension associated events from five short-term, placebo-controlled trials in schizophrenia (n=926) on ABILIFY included: orthostatic hypotension (placebo 1%, aripiprazole 1.9%), orthostatic lightheadedness (placebo 1%, aripiprazole 0.9%), and syncope (placebo 1%, aripiprazole 1.9%), orthostatic lightheadedness sosciated events from short-term, placebo-controlled trials in bipolar mania (n=597) on ABILIFY included: orthostatic hypotension (placebo 0%, aripiprazole 0.7%), orthostatic lightheadedness (placebo 1.5%, aripiprazole 0.5%), and syn-cope (placebo 1%, aripiprazole 0.5%), the incidence of a significant orthostatic charget in blood pressure (defined as a decrease of at least 30 mmHg in systolic blood pressure when charging from a supine to standing position) for aripiprazole was not statistically different from placebo (in exclosphrenia: 14% among aripiprazole-treated patients and 12% among placebo-treated patients). Aripiprazole should be used with caution in patients with known cardiovascular disease history of movardial infarction or ischemic heart disease. heart failure or conduction abnormalifies), cerebrovascular history of myocardial infarction or ischemic heart disease, heart failure or conduction abnormalities), cerebrovascular disease, or conditions which would predispose patients to hypotension (dehydration, hypovolemia, and treatment with antihypertensive medications). Seizure

Seizure Soccurred in 0.1% (1/926) of aripiprazole-treated patients with schizophrenia in short-term, placebo-controlled trials. In short-term, placebo-controlled clinical trials of patients with bipolar mania, 0.3% (2/597) of aripiprazole-treated patients and 0.2% (1/436) placebo-treated patients experienced seizures. As with other antipsychotic drugs, aripiprazole should be used cautiously in patients with a history of seizures or with conditions that lower the seizure threshold, e.g., Alzheimer's dementia. Conditions that lower the seizure threshold may be more prevalent in a population of 65 years or older.

Potential for Cognitive and Motor Impairment

in short-term, placebo-controlled trials of schizophrenia, somolence was reported in 11% of patients on ABILIFY com-pared to 8% of patients on placebo; somolence led to discontinuation in 0.1% (1/926) of patients with schizophrenia on ABILIFY in short-term, placebo-controlled trials. In short-term, placebo-controlled trials of blookar mania, somnolence was reported in 14% of patients on ABILIFY compared to 7% of patients on placebo, but id in on lead to discontinuation of any patients with bipolar mania. Despite the relatively modest increased incidence of somnolence compared to placebo, Patients with operating bacardous machinery, including automobiles, until they are reasonably certain that therapy with ABILIFY like other antipsycholos, may have the potential to impair judgment, thinking, or motor skills. Patients should be cautioned about operating hazardous machinery, including automobiles, until they are reasonably certain that therapy with ABILIFY does not affect them adversely.

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Body Temperature Regulation

Disruption of the body's ability to reduce core body temperature has been attributed to antipsychotic agents. Appropriate Disciplication in the body is during to reduce core body reimpenditie has obten autorout or antibyto fold eghts. Appropriate care is advised when prescribing aripiprazole for patients who will be experiencing conditions which may contribute to an elevation in core body temperature, e.g., exercising strenuously, exposure to extreme heat, receiving concomitant medication with anticholinergic activity, or being subject to dehydration. Dvsphaqia

Ecophageal dysmotility and aspiration have been associated with antipsychotic drug use. Aspiration pneumonia is a com-mon cause of morbidity and mortality in elderly patients, in particular those with advanced Alzheimer's dementia. Arlpiprazole and other antipsychotic drugs should be used cautiously in patients at risk for aspiration pneumonia (see PRECAUTIONS: Use in Patients with Concomitant Illness). Suicide

The possibility of a suicide attempt is inherent in psychotic illnesses and bipolar disorder, and close supervision of high-risk patients should accompany drug therapy. Prescriptions for ABILIFY (aripiprazole) should be written for the smallest quantity of tablets consistent with good patient management in order to reduce the risk of overdose. Use in Patients with Concomitant Illness

Safety Experience in Elderly Patients with Psychosis Associated with Alzheimer's Disease: In a flexible dose (2 to Safety experience in cuery returns with responses associated with Alzheime's Disease in a instole dose (2 to 15 mg/day). To week, placebo-controlled study of aripiprozole in elderty platents (mean age: 81.5 years; ange: 56 to 95 years) with psychosis associated with Alzheime's dementia, 4 of 105 patients (3.8%) who received ABILIFY died com-pared to no deaths among 102 patients who received placebo during or within 30 days after termination of the double-blind portion of the study. Three of the patients (age 92, 91, and 87 years) died following the discontinuation of ABILIFY in the double-blind phase of the study (causes of death were pneumonia, heart failure, and shock). The fourth patient (age The boole company present of the study (classes of une study) (classes of une study). The study. The treat ment-emergent adverse events that were reported at an incidence of ≥5% and having a greater incidence than placebo in this study were acci-dental injury, somnolence, and bronchitis. Eight percent of the ABILIFY-treated patients reported somnolence compared to one percent of placebo patients. In a small plicit, open-label, ascending-dose, cohort study (n=30) in eldely patients. to one percent of placebo patients. In a small piot, open-label, ascending-dose, cohort study (in=30) in elderly patients with dementia, ABILFY was associated in a dose-related tashion with somolence. The safety and efficacy of ABILFY in the treatment of patients with psychosis associated with dementia have not been established. If the prescriber elects to treat such patients with ABILFY wigilance should be exercised, particularly for the emergence of difficulty swallowing or excessive somolence, which could predispose to accidental injury or aspiration. Clinical experience with ABILFY in patients with certain concomitant systemic illnesses (see CLINICAL PHARIMACOLOGY. Special Populations: Renal Impairment and Hepatic Impairment in Full Prescribing Information) is limited. ABILFY has not been evaluated or used to any appreciable extent in patients with a recert history of myocardial infarction or unstable heart disease. Patients with these diagnoses were excluded from premarketing clinical studies.

Information for Patients: Physicians are advised to consult full prescribing information to review issues to be discussed wtih patients for whom they prescribe ABILIFY.

Drug-Drug Interactions: Given the primary CNS effects of aripiprazole, caution should be used when ABILIFY is taken in brug Proving interactions: twen the primary two enects of anophrazole, caution should be used when hour's taken in combination with other centrally acting drugs and alcohol. Due to its or, adrenergic receptor antagonism, anipprazole has the potential to enhance the effect of certain anthypertensive agents. *Potential for Other Drugs to Affect ABL/PY*: Aripiprazole is not a substrate of CYP1A1, CYP1A2, CYP2A6, CYP2C6, CYP2C6, CYP2C19, or CYP2E1 enzymes. Aripiprazole also does not undergo direct glucuronidation. This suggests that an interaction of antipiprazole with inhibitors Anpiprazole is not a substrate of CYP1A1, CYP1A2, CYP2A6, CYP2C9, CYP2 architecture uses withdrawn from the combination therapy, anpiprazio should be bauld the formed valuation of significant effect of famotidine, valproate, or lithium was seen on the pharmacokinetics of anpipraziole (see CLINICAL PHARMACOLOGY: Drug-Drug Interactions in Full Prescribing Information). Potential for ABIL/FY to Affect Other Drugs: Aripiprazole is unlikely to cause clinically important pharmacokinetic interactions with drugs metabolized by sytochrome Application is universe in a way obtained in the analysis of a provide the advectory of a state of the advectory of the advec motor skills or stimulus response in healthy subjects. As with most psychoactive medications, patients should be advised to avoid alcohol while taking ABILIFY.

Carcinogenesis, Mutagenesis, Impairment of Fertility: (Please See Full Prescribing Information).

Pregnancy

Pregnancy Pregnancy Category C In animal studies, aripiprazole demonstrated developmental toxicity, including possible teratogenic effects in rats and rabbits. Pregnant rats were treated with oral doses of 3, 10, and 30 mg/kg/day (1, 3, and 10 times the maximum rec-ommended human dose [MRHD] on a mg/m² basis) of aripiprazole during the period of organogenesis. Gestation was the uncertained and the Tradmod towards a slight delay in fatal downlowment as evidenced by decreased fetal ommended human dose [MRHD] on a mg/m⁵ basis) of aripiprazole during the period of organogenesis. Gestation was slightly prohonged at 30 mg/kg. Treatment caused a slight telday in fetal development, as evidenced by decreased fetal weight (30 mg/kg), undescended testes (30 mg/kg), and delayed skeletal ossification (10 and 30 mg/kg). There were no adverse effects on embryofetal or pup survival. Delivered offspring had decreased bodyweights (10 and 30 mg/kg), and increased incidences of hepatodiaphragmatic nodules and diaphragmatic hemia at 30 mg/kg (the other dose groups were not examined for these findings). (A low incidence of diaphragmatic hemia was also seen in the fetuses exposed to 30 mg/kg.) Postnatily, delayed vaginal opening was seen at 10 and 30 mg/kg and imgraid reproductive performance (decreased fertility rate, corpora lutea, implants, and live fetuses, and increased post-implantation loss, likely mediated (decreased fertility rate, corpora lutea, implants, and live fetuses, and increased post-implantation loss, likely mediated through effects on female offspring) was seen at 30 mg/kg, Some maternal toxicity was seen at 30 mg/kg, however, there was no evidence to suggest that these developmental effects were secondary to maternal toxicity. Pregnant rabibits were treated with oral doses of 10, 30, and 100 mg/kg/day (2, 3, and 11 times human exposure at MRHo based on AUC and 6, 19, and 65 times the MRHD based on mg/m³) of aripiprazole during the period of organogenesis. Decreased maternal food consumption and increased abortions were seen at 100 mg/kg, treatment caused increased fetal weight (30 and 100 mg/kg), increased incidence of skeletal abnormality (fused sternebrae at 30 and 100 mg/kg), and minor skeletal variations (100 mg/kg). In a study in which rats were treated with oral doses of 3, 10, and 30 mg/kg/day (1, 3, and 10 times the MRHD on a mg/m² pasis) of anipiprazole perinatally and posthatally (from day 17 of gestation through day 21 postpartum), slight maternal toxicity and slightly prolonged gestation were seen at 30 mg/kg, an increase in stillbirths, and decreases in pup weight (persisting into adulthood) and surival, were seen at 30 mg/kg. at this dose. There are no adequate and well-controlled studies in pregnant women. It is not known whether aripiprazole can cause fetal harm when administered to a pregnant woman or can affect reproductive capacity. Aripiprazole should be used during pregnancy only if the potential benefit outweighs the potential risk to the fetus. Labor and Delivery

The effect of aripinrazole on labor and delivery in humans is unknown.

Nursing Mothers

Aripiprazole was excreted in milk of rats during lactation. It is not known whether aripiprazole or its metabolites are excreted in human milk. It is recommended that women receiving aripiprazole should not breast-feed. Pediatric Use

Safety and effectiveness in pediatric and adolescent patients have not been established.

Geriatric Use

Of the 7551 patients treated with aripiprazole in premarketing clinical trials, 991 (12%) were ≥65 years old and 789 (10%) were ≥75 years old. The majority (88%) of the 991 patients were diagnosed with dementia of the Alzheimer's type. (10%) were ≥75 years old. The majorfty (86%) of the 991 patients were diagnosed with dementia of the Alzheimer's type. Placebo-controlled studies of aripiprazole in schizophrenia or bipolar mania did not include sufficient numbers of sub-jects aged 65 and over to determine whether they respond differently from younger subjects. There was no effect of age on the pharmacokinetics of a single 15-mg dose of aripiprazole. Aripiprazole clearance was decreased by 20% in elder-ly subjects (≥65 years) compared to younger adult subjects (18 to 64 years), but there was no detectable effect of age in the population pharmacokinetic analysis in schizophrenia patients. Studies of elderly patients with psychosis associ-ated with Alzheimer's disease have suggested that there may be a different tolerability profile in this population pared to younger patients with schizophrenia (see **PRECAUTIONS**: Use in *Patients with Concomitant lilness*). The safety and efficacy of ABLIFY in the treatment of patients with psychosis associated with Alzheimer's disease has no been established. If the prescriber elects to treat such patients with ABLIFY, vigilance should be exercised.

ADVERSE REACTIONS

Aripiprazole has been evaluated for safety in 7951 patients who participated in multiple-dose, premarketing trials in schizophrenia, bipolar mania, and dementia of the Alzheimer's type, and who had approximately 5235 patient-years of exposure. A total of 2280 aripiprazole-treated patients were treated for at least 180 days and 1558 aripiprazole-treated patients had at least 1 year of exposure. The conditions and duration of treatment with anipiprazole included (in overlap-ping categories) double-blind, comparative and noncomparative open-label studies, inpatient and outpatient studies ping categories double-bind, comparative and noncomparative open-haloe soules, imparent and outparent soules, fixed- and flexible-does studies, and short- and longer-term exposure. Adverse events during exposure were obtained by collecting volunteered adverse events, as well as results of physical examinations, vital signs, weights, laboratory analyses, and ECG. Adverse experiences were recorded by clinical investigators using terminology of their own choosing. In the tables and tabulations that follow, modified COSTANT dictionary terminology and been used initially to classify reported adverse events into a smaller number of standardized event categories, in order to provide a meaningful estireported adverse events into a smaller intrinser or standardized event categories, in order to provide a meaningfue sai-mate of the proportion of individuals experiencing adverse events. The stated frequencies of adverse events represent the proportion of individuals who experienced at least once, a treatment-emergent adverse events represent was considered treatment emergent if it occurred for the first time or worsened while receiving therapy following baseline evaluation. There was no attempt to use investigator causality assessments; i.e., all reported events are includ-ed. The prescriber should be aware that the figures in the tables and tabulations cannot be used to predict the incidence etc. The prescriber should be aware that the righters in the tables and tabulators calmot be used to provide the income of side effects in the course of usual medical practice where patient characteristics and other factors differ from those that prevailed in the clinical trials. Similarly, the cited frequencies cannot be compared with figures obtained from other clinical investigations involving different treatment, uses, and investigators. The cited figures, however, do provide the prescribing physician with some basis for estimating the relative contribution of drug and nondrug factors to the adverse event incidence in the population studied.

event incidence in the population studied. Adverse Findings Observed in Short-Term, Placebo-Controlled Trials of Patients with Schizophrenia The following findings are based on a pool of five placebo-controlled trials (four 4-week and one 6-week) in which arigiprazole was administered in doese ranging from 2 to 30 mg/day. Adverse Events Associated with Discontinuation of Treatment in Short-Term, Placebo-Controlled Trials Overall, there was no difference in the incidence of discontinuation due to adverse events between aripiprazole-treated

(7%) and placebo-treated (9%) patients. The types of adverse events that led to discontinuation were similar between the aripiprazole and placebo-treated patients.

amprizzore and pracebo-treated patients. Adverse Findings Observed in Short-Term, Placebo-Controlled Trials of Patients with Bipolar Mania The following findings are based on a pool of 3-week, placebo-controlled bipolar mania trials in which aripiprazole was administered at doses of 15 or 30 mg/day. Adverse Events Associated with Discontinuation of Treatment in Short-Term, Placebo-Controlled Trials Overall, in patients with bipolar mania, there was no difference in the incidence of discontinuation due to adverse events Uverai, in patients with oppoar mania, there was no omerence in the incidence of obscontinuation out to adverse events between anjoinzaole-treated (11%) and placebo-treated (9%) patients. The types of adverse events that led to discon-tinuation were similar between the aripiprazole and placebo-treated patients. Commonly Observed Adverse Events in Short-Term, Placebo-Controlled Trials of Patients with Bipolar Mania Commonly Observed Adverse events associated with the use of aripiprazole in patients with bipolar mania (incidence of 5% or greater and aripiprazole incidence at least twice that for placebo) are shown in Table 1. There were no adverse

events in the short-term trials of schizophrenia that met these criteria.

Table 1: Commonly Observed Adverse Events in Short-Term, Placebo-Controlled Trials in Patients with Bipolar Mania

	Percentage of Patients Reporting Event		
Adverse Event	Aripiprazole (n=597)	Placebo (n=436)	
Accidental Injury	6	3	
Constipation	13	6	
Akathisia	15	4	

Adverse Events Occurring at an incidence of 2% or More Among Aripiprazole-Treated Patients and Greater than Placebo in Short-Term, Placebo-Controlled Trials

Treatment-emergent adverse events that occurred during acute therapy (up to 6 weeks in schizophrenia and up to 3 weeks in biologram mania), including only those events that occurred in 2% or more of patients treated with aripiprazole (doses ≥2 mg/day) and for which the incidence in patients treated with aripiprazole was greater than the incidence in (uoses 22 mg/oay) and or which the incidence in patients treated with apprature was greater train the incidence in patients treated with placebox in the combined dataset were. Body as a Whole—headache, asthenia, accidental injury, peripheral edema; Cardiovascular System—hypertension; Digestive System—nausea, dyspepsia, vomiting, constipation; Muscubskietal System—myaigia, Nervous System—egitation, anxiety, insormia, somnolence, akathenia, acidentia, injurg, Special Senses—blurred vision. An examination of population subgroups did not reveat any clear evidence of differential adverse event incidence on the basis of age, gender, or race.

Dose-Related Adverse Events

Schizophrenia

Does response relationships for the incidence of treatment-emergent adverse events were evaluated from four trials in patients with schizophrenia comparing various fixed doses (2, 10, 15, 20, and 30 mg/day) of aripiprazole to placebo. This analysis, stratified by study, indicated that the only adverse event to have a possible dose response relationship, and then most prominent only with 30 mg, was sommalence (placebo, 7.7%, 15 mg, 8.7%, 20 mg, 7.5%, 30 mg, 15.3%). Extrapyramidal Symptoms In the short-term, placebo-controlled trials of schizophrenia, the incidence of reported EPS for aripiprazole-treated patients

In the short-term, placebo-controlled unais of schizophrenia, the incidence of reported ErS for an plazacio-reated plauents was 6% us. 6% for placebo. In the short-term, placebo-controlled trials in bipolar mania, the incidence of reported ErS-related events excluding events related to akathisia for aripiprazole-treated patients was 17% vs. 12% for placebo. In the short-term, placebo-controlled trials in bipolar mania, the incidence of akathisia-related events for aripiprazole-treated patients was 15% vs. 4% for placebo. Dipolar mania, the incidence of akathisia-related events for aripiprazole-treated patients was 15% vs. 4% for placebo. Dipolar mania, the incidence of akathisia-plated the selected on the Simpson Angus Rating Scale (for EPS), the Barnes Akathisia Scale (for akathisia) and the Assessments of Involuntary Movement Scales Rating Scale (for EPS), the Barnes Akathisia Scale (for akathisia) and the Assessments of Involuntary Movement Scales (for dyskinesias). In the schizophrenia trials, the objectively collected data did not show a difference between ania trials, the Simpson Angus Rating Scale and the Barnes Akathisia Scale (aripiprazole, 0.08; placebo, -0.05), in the bipolar mania trials, the Simpson Angus Rating Scale and the Barnes Akathisia Scale showed a significant difference between aripiprazole and placebo (aripiprazole), 0.61; placebo, 0.03 and aripiprazole, 0.25; placebo, -0.06). Changes in the Assessments of Involuntary Movement Scales were similar for the aripiprazole, 0.25; placebo, -0.06). Changes in the Assessments of Involuntary Movement Scales were similar for the aripiprazole, 0.25; placebo, -0.06). Changes in the (for EPS), the Barnes Akathisia Scale (for akathisia) and the Assessments of Involuntary Movement Scales (for dyskinesias) did on choru a difference between aripiprazole and placebu and the Assessments of Involuntary Movement Scales (for dyskinesias). did not show a difference between aripiprazole and placebo. Laboratory Test Abnormalities

Laboratory lest Anormalities A between group comparison for 3- to 6-week, placebo-controlled trials revealed no medically important differences between the aripiprazole and placebo groups in the proportions of patients experiencing potentially clinically significant changes in routine serum chemistry, hematology, or urinalysis parameters. Similarly, there were no aripiprazole/placebo differences in the incidence of discontinuations for changes in serum chemistry, hematology, or urinalysis. In a long-term (26-week), placebo-controlled trial there were no medically important differences between the aripiprazole and placebo patients in the mean change from baseline in prolactin, fasting glucose, triglyceride, HDL, LDL, and total cholesterol measurements.

Weight Gain

Weight Cain In 4- to 6-week trials in schizophrenia, there was a slight difference in mean weight gain between aripiprazole and placebo patients (+0.7 kg vs. -0.05 kg, respectively), and also a difference in the proportion of patients meeting a weight gain criterion of \geq 7% of body weight [anipiprazole (8%) compared to placebo (3%)]. In 3-week trials in mania, the mean weight gain for aripiprazole and placebo patients was 0.0 kg vs. -0.2 kg, respectively. The proportion of patients meet-ing a weight gain criterion of \geq 7% of body weight was aripiprazole (3%) compared to placebo (2%). Table 2 provides the weight change results from a long-term (26-week), placebo-controlled study of aripiprazole, both mean change from baseline and proportions of patients meeting a weight gain criterion of \geq 7% of body weight relative to baseline, catego-rized by BMI at baseline:

Table 2: Weight Change Results Categorized by BMI at Baseline: Placebo-Controlled Study in Schizophrenia, Safety Sample						
	BMI <23		BN	1 23-27	BN	AI >27
	Placebo	Aripiprazole	Placebo	Aripiprazole	Placebo	Aripiprazole
Mean change from baseline (kg)	-0.5	-0.5	-0.6	-1.3	-1.5	-2.1
% with >7% increase BW	3.7%	6.8%	4 2%	5 1%	41%	5 7%

Table 3 provides the weight change results from a long-term (52-week) study of aripiprazole, both mean change from ence o provides are weight charge results from a long-term (s2-week) study of anjpiprazole, both mean change from baseline and proportions of patients meeting a weight gain criterion of ≥7% of body weight relative to baseline, catego-rized by BM at baseline:

Table 3: Weight Change Results Categorized by BMI at Baseline: Active-Controlled Study in Schizophrenia, Safety Sample					
	BMI <23	BMI 23-27	BMI >27		
Mean change from baseline (kg)	2.6	1.4	-1.2		
% with >7% increase BW	30%	19%	8%		

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ECG Changes

Between group comparisons for pooled, placebo-controlled trials in patients with schizophrenia, revealed no significant differences between aripiprazed and placebo in the proportion of patients experiencing potentially important changes in ECG parameters, in fact, within the dose rance of 10 to 30 molday, anipiprazele tended to silohtly shorten the 0T, interval. Aripiprazole was associated with a median increase in heart rate of 4 beats per minute compared to a 1 beat per minute increase among placebo patients. Additional Findings Observed in Clinical Trials

Adverse Events in a Long-Term, Double-Blind, Placebo-Controlled Trial The adverse events reported in a 26-week, double-blind trial comparing ABILIFY (aripiprazole) and placebo were generally consistent with those reported in the short-term, placebo-controlled trials, except for a higher incidence of tremor [9% (13/153) for ABILIFY vs. 1% (2/153) for placebo]. In this study, the majority of the cases of tremor were of mild (9% (13/15)) for AbiLit Ys. 1% (2/15) for placeop) in this study, the majority of the cases or leminow were of limited duration (9/13 ≤10 days). Tremor infrequently led to discontinuation (<1%) of ABILIFY. In addition, in a long-term (52-week), active-controlled study, the incidence of tremor for ABILIFY was 4% (34/859). **Other Adverse Events Observed During the Premarketing Evaluation of Aripiprazole** Following is a list of modified COSTART terms that reflect treatment-emergent adverse events as defined in the intro-

duction to the ADVERSE REACTIONS section reported by patients treated with aripiprazole at multiple doses >2 mg/day during any phase of a trial within the database of 7951 patients. All reported events are included except those already listed in other parts of the ADVERSE REACTIONS section, those considered in the WARNINGS or PRECAUTIONS, those event terms which were so general as to be uninformative, events reported with an incidence of <0.05% and which did not have a substantial probability of being acutely life-threatening, events that are otherwise common as background events, and events considered unlikely to be drug related. It is important to emphasize that, although the events reportevents, and events considered uniter vibre of due to the standard of the standard of the events are further categorized by ed occurred during treatment with arbiprozel, they were not necessarily caused by it. Events are further categorized by body system and listed in order of decreasing frequency according to the following definitions: frequent adverse events are those occurring in at least 1/100 patients (only those not already listed in the tabulated results from placebo-con-trolled trials appear in this listing); infrequent adverse events are those occurring in 1/100 to 1/1000 patients; are events are those occurring in fewer than 1/1000 patients. *Body as a Whole: Frequent* – flu syndrome, fever, clest pain, rigidity including neck and extremity), neck pain, pelvic pain, *infraquent* – face defma, suicide attment, mana painter, chills, photosensitivity, tightness (including addomen, back, extremity, head, jaw, neck, and tongue), jaw pain, bloating, enlarged addomen, chest tightness, throat pain; *Rare* – moniliasis, head heaviness, throat tightness, therde pain; *Rare* – moniliasis, head heaviness, throat tightness, therden pain; *Rare* – tachycardia (including ventricular and supraventricular), hypotension, bradycardia: Infrequent – palpitation, hemorrhage, heart failure, myocardial infarction, cardiac arrest, atrial fibrillation, AV block, prolonged QT interval, extrasystoles, myocardial ischemia, deep vein thrombosis, angina pectoris, pallor, cardiopulmonary arest, philoitiis, Rare – bundle branch block, atrial futter, vasovagi reaction, cardiomegaly, thrombophilebitis, car-diopulmonary failure. *Digestive System: Frequent* – nausea and vomiting; *Intrequent* – increased appetite, dysphagia, gastroenteritis, flatulence, tooth caries, gastritis, gingivitis, gastrointestinal hemorrhage, hemorrhoids, gastroesophageal reflux, periodontal abscess, fecal incontinence, rectal hemorrhage, stomatitis, colitis, tongue edema, cholecystitis, mouth ulcer, oral moniliasis, eructation, fecal impaction, cholelithiasis; Rare – esophagitis, hematemesis, intestinal obstruction, gum hemorrhage, hepatitis, peptic ulcer, glossitis, melena, duodenal ulcer, cheilitis, hepatomegaly, pancreatitis. Endocrine System: Infrequent – hypothyroidism; Rare – goiter, hyperthyroidism. Hemic/Lymphatic System: Frequent – ecchymosis, anemia; infrequent – hypochromic anemia, leukocytosis, leukopenia (including neutropenia), lymphadenopathy, eosinophilla, macrocytic anemia; *Rare* – thrombocythemia, thrombocytopenia, petechlae. *Metabolic and Nutritional Disorders: Frequent* – weight loss, creatine phosphokinase increased, dehydration; *Infrequent* – edema, hyperglycemia, hypercholesteremia, hypokalemia, diabetes mellitus, hypoglycemia, hyperilperina, SGPT increased, thirst, BUN increased, hyponatremia, SGOT increased, creatinine increased, cyanosis, alkaline phosphatase increased, biinubinemia, iron defi-ciency anemia, hyperkalemia, hyperuricemia, obesity; *Rare* – lactic dehydrogenase increased, hypernatremia, gout, hypo-glycemic reaction. *Musculoskeletal System: Frequent* – muscle cramp; *Infrequent* – arthralgia, myasthenia, arthrosis, bone pain, arthritis, muscle weakness, spasm, bursitis, myopathy; *Rare* – rheumatoid arthritis, rhabdomyolysis, ten-donitis, tenosynovitis. *Nervous System: Frequent* – depression, nervousness, schizophrenic reaction, hallucination, hostility, confusion, paranoid reaction, suicidal thought, abnormal gait, manic reaction, delusions, abnormal dream; Infrequent – emotional lability, twitch, cogwheel rigidity, impaired concentration, dystonia, vasodilation, paresthesia, impotence, extremity tremor, hypesthesia, vertigo, stupor, bradykinesia, apathy, panic attack, decreased libido, hypersonnia, dyski-nesia, manic depressive reaction, ataxia, visual hallucination, cerebrovascular accident, hypokinesia, depersonalization, impaired memory, delinium, dysarthria, tardive dyskinesia, amnesia, hyperactivity, increased libido, myocionus, restless leg, neuropathy, dysphoria, hyperkinesia, cerebral ischemia, increased reflexes, akinesia, decreased consciousness, hyperesthesia, slowed thinking; Rare - blunted affect, euphoria, incoordination, oculogyric crisis, obsessive thought, hypotonia, buccoglossal syndrome, decreased reflexes, derealization, intracranial hemorrhage. Respiratory System: Inprotons, bacograduate of market and a standard and a standard and a standard and a standard of postandard of postandard of postandard and a standard of postandard and a standard and nemocysis contrain population, alopecal seborriea, protaisis, Rare – maculopapular rash, exfoliative dermatitis, arcne, eczema, skin discoloration, alopecal, seborriea, psoriasis, Rare – maculopapular rash, exfoliative dermatitis, urticaria. Special Senses: Frequent – conjunctivitis, Infrequent – ear pain, dry eye, eye pain, tinnitus, cataract, otilis media, altered taste, blepharitis, eye hemorrhage, deafness; Rare – diplopia, frequent blinking, ptosis, otilis externa, ambigopia, photophobia. Urogenital System: Frequent – urinary incontinence; Infrequent – urinary frequency, leukorrhea, amblyopia, photophobia. Urogenital System: Frequent – urinary incomtinence; Infrequent – urinary requency, leukorrhea, urinary retention, cystitis, hematuria, dysuria, amenorrhea, vaginal hemorrhage, abnormal ejaculation, kidney failure, vaginal monilissis, urinary urgency, gynecomastia, kidney calculus, albuminuria, breast pain, urinary burnling; Rare – noc-turia, polyuria, menorrhagia, anorgasmy, glycosuria, cervicitis, uterus hemorrhage, female lactation, urolithiasis, priapism. **Other Events Observed During the Postmarketing Evaluation of Aripiprazole** Voluntary reports of adverse events in patients laking aripiprazole that have been received since market introduction and not listed above that may have no causal relationship with the drug include rare occurrences of allergic reaction (e.g., anaphylactic reaction, angioedema, laryngospasm, pruritis, or urticaria).

DRUG ABUSE AND DEPENDENCE

Controlled Substance

ABILIFY is not a controlled substance.

Abuse and Dependence

Aripiprazole has not been systematically studied in humans for its potential for abuse, tolerance, or physical dependence. In physical dependence studies in monkeys, withdrawal symptoms were observed upon abrupt cessation of dosing. While the clinical trials did not reveal any tendency for any drug-seeking behavior, these observations were not systematic and it is not possible to predict on the basis of this limited experience the extent to which a CNS-active drug will be misused, diverted, and/or abused once marketed. Consequently, patients should be evaluated carefully for a history of drug abuse, and such patients should be observed closely for signs of ABILIFY misuse or abuse (e.g., development of tolerance, increases in dose, drug-seeking behavior).

OVERDOSAGE Management of Overdosage

No specific information is available on the treatment of overdose with aripiprazole. An electrocardiogram should be obtained in case of overdosage and, if QT, interval prolongation is present, cardiac monitoring should be instituted. Otherwise, management of overdose should concentrate on supportive therapy, maintaining an adequate airway, oxygenation and ventilation, and management of symptoms. Close medical supervision and monitoring should continue until the patient recovers.

Charcoal: In the event of an overdose of ABILIFY, an early charcoal administration may be useful in partially prevent-ing the absorption of aripiprazole. Administration of 50 g of activated charcoal, one hour after a single 15-mg oral dose aripiprazole, decreased the mean AUC and C_{max} of aripiprazole by 50%. Hemodialysis: Although there is no information on the effect of hemodialysis in treating an overdose with aripiprazole,

hemodialysis is unlikely to be useful in overdose management since aripiprazole is highly bound to plasma proteins. Storage

Store at 25° C (77° F); excursions permitted to 15-30° C (59-86° F) [see USP Controlled Room Temperature]. Marketed by Otsuka America Pharmaceutical, Inc., Rockville, MD 20850 USA

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🚱 Otsuka America Pharmaceutical, Inc. Rockville, MD 20850 U.S.A.

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