Volume 3 - Number 3

CNS SPECTRUMS

## The International Journal of Neuropsychiatric Medicine

## Autism: Emerging Consensus

A Dimensional Approach to the Autism Spectrum E. Hollander Autism Screening and Diagnostic Evaluation: CAN Consensus Statement CAN Consensus Group Progress in the Neurobiology of Autism I. Rapin Seizures and EEG Findings in Children With Autism Spectrum Disorder R. Tuchman

An Immunologic Theory for the Development of Some Cases of Autism R. P. Warren Autism, Serotonin, and the Cerebellum: Might There Be a Connection? D. Marazziti

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**Photo Essay** Autism—one of the few neuropsychiatric illnesses to strike in early childhood—is depicted by the universal image of a young person in a state of social isolation. This issue, dedicated to an emerging consensus in autism, includes first-time presentations in the field. **Articles Inside**.

# Once-a-day ARICEPT® (donepezil HCl)– First-line therapy for mild to moderate Alzheimer's disease

## PROVEN EFFECTIVE IN ENHANCING COGNITIVE FUNCTION

Effect on cognitive function over 24 weeks of active treatment and 6 weeks of placebo as measured by ADAS- $cog^{1*}$ 



<sup>e</sup>Alzheimer's Disease Assessment Scale, cognitive subscale (ADAS-cog) is a 70-point, clinically validated psychometric scale for measuring cognitive function in patients with Alzheimer's disease. In one controlled clinical trial of 30 weeks' duration in 473 patients, 154 patients were randomly assigned to receive daily doses of 5 mg. One hundred fifty-seven patients were randomly assigned to receive daily doses of 5 mg. One hundred trial of 30-week trial was divided into a 24-week double-blind active treatment phase followed by a 6-week single-blind placebo washout period.

- Significant benefits observed in 24-week study in both 5 mg/day and 10 mg/day ARICEPT<sup>®</sup> groups
- Placebo washout demonstrates that beneficial effects of ARICEPT<sup>®</sup> abate following discontinuation

Please see brief summary of prescribing information on the last page of this advertisement.

Reference: I. Rogers SL, Farlow MR, Doody RS, et al. A 24-week, double-blind, placebo-controlled trial of donepezil in patients with Alzheimer's disease. *Neurology*. 1998;50:136-145. ARICEPT® is a registered trademark of Eisai Co., Ltd.

## EXPERIENCE & CONVENIENCE

- Over 250,000 prescriptions written to date
- Once-daily administration, with or without food
- Some patients might derive additional benefit from escalation to 10-mg daily after 4 to 6 weeks of 5-mg once-daily therapy

## SAFETY & TOLERABILITY

- No liver function testing required
- No significant drug-drug interactions observed in clinical trials with the following commonly prescribed medications: cimetidine, digoxin, theophylline, and warfarin
- The most common adverse events leading to discontinuation in clinical trials with ARICEPT<sup>®</sup> were nausea, diarrhea, and vomiting
- Clinical studies of ARICEPT<sup>®</sup> have shown no increase, relative to placebo, in the incidence of either peptic ulcer disease or gastrointestinal bleeding. Nevertheless, cholinesterase inhibitors may be expected to increase gastric acid secretion. Therefore, patients (especially those at increased risk for developing ulcers — eg, history of ulcer disease, receiving concurrent nonsteroidal anti-inflammatory drugs) should be monitored closely for gastrointestinal bleeding
- In clinical trials, syncopal episodes have been reported in association with the use of ARICEPT<sup>®</sup> (2% vs 1% for placebo)





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## ARICEPT (donepezil HCl)

THERAPY TO REMEMBER

#### ARICEPT<sup>®</sup> (Donepezil Hydrochloride Tablets)

ARICET<sup>1</sup> Conference in the second provide interest of the second provided and succinvictoriline-type muscle relaxation during anesthesia. **Cardiovascular Conditions:** Because of their pharmacological action, cholinesterase inhibitors may have vagotonic effects on heart rate (eg, bradycardia). The potential for this action may be particularly important to patients with "sick sinus syndrome" or other supraventricular cardiac conduction conditions. Syncopal episodes have been reported in association with the use of ARICEPT<sup>e</sup>. **Gastriointestinal Conditions:** Through their primary action, cholinesterase inhibitors may be expected to increase gastric acid secretion due to increased cholinergic activity. Therefore, patients should be monitored closely for symptoms of active or occult gastrointestinal bleeding, especially those at increased risk for developing ulcers, eq. those with a history of ulcer disease or those receiving concurrent nonsteroidal anti-inflammatory drugs (NSAIDS) Clinical studies of ARICEPT® have shown no increase, relative to placebo, in the incidence of either peptic ulcer Clinical studies of AHICEPT have shown no increase, relative to placebo, in the incidence of either peptic ulcer disease or gastrointestinal bleeding. ARICEPT<sup>®</sup>, as a predictable consequence of its pharmacological properties, has been shown to produce diarrhea, nausea, and vomiting. These effects, when they occur, appear more frequently with the 10 mg/day does than with the 5 mg/day does. In most cases, these effects have been mild and transient, sometimes lasting one to three weeks, and have resolved during continued use of ARICEPT<sup>®</sup>. Gentinourinary: Although not observed in clinical trials of ARICEPT<sup>®</sup>, cholinomimetics may cause bladder outflow obstruction. **Neurological Conditions:** Selzures: Cholinomimetics are believed to have some potential to cause generalized convulsions. **However**, selzure activity also may be a mainficiation of Atheriner's Disease. **Purimoary Conditions:** Because of their cholinomimetic actions, cholinesterase inhibitors should be prescribed with care to patients with a history of them or pharmed disease disease. their cholinomimetic actions, cholinesterase inhibitors should be prescribed with care to patients with a history of asthma or obstructive pulmonary disease. **PRECAUTIONS Drug-Drug Interactions Drugs Highly Bound to Plasme Proteins:** Drug displacement studies have been performed *in vitro* between this highly bound drug (96%) and other drugs such as furosemide, dipoxin, and warfarin. ARICEPT\* at concentrations of 0.3-10 µg/mL did not affect the binding of turosemide (5 µg/mL), digoxin (2 ng/mL), and warfarin (3 µg/mL) to human albumin. Similarly, the binding of ARICEPT\* to human albumin was not affected by furosemide, digoxin and warfarin. **Effect of ARICEPT\* on the Metabolism of Other Drugs:** No *in vivo* clinical trials have investigated the effect of **ARICEPT\* on the detactors** of drugs metabolized by CYP 3A4 (eq. cisapride, terfenadine) or by CYP 2D6 (eq. imigrarnine). However, *in vitro* studies show a low rate of binding to these enzymes (mean K, about 50 - 130 µM), that, given the therapeutic plasma concentrations of donepezil (164 nM), indicates little likelihood of interference. Whether **ARICEPT\*** has any potential for enzyme induction is not known. Formal pharmacokinetic studies evaluated the potential **ABICEPT\*** for interactions with the pharking curvative quarkaria and dioxin. No significant effects on the **ABICEPT\*** has any potential for enzyme induction is not known. Formal pharmacokinetic studies evaluated the potential **ABICEPT\*** has any potential for enzyme induction is not known. Formal pharmacokinetic studies evaluated the potential **ABICEPT\*** has any potential for enzyme induction is not known. Formal pharmacokinetic studies evaluated the potential **ABICEPT\*** has any potential for enzyme induction is not known. Formal pharmacokinetic studies evaluated the potential **ABICEPT\*** has any potential for enzyme induction is not known. Formal pharmacokinetic studies evaluated the potential **ABICEPT\*** has any potential for enzyme induction is not known. Formal pharmacokinetic s AndCFT task and polania for each induction that with a non-induction polarization such sevena and the polarization of ARICEFT<sup>®</sup> for interaction with the ophylline, cimetidine, warfarin and digoxin. No significant effects on the pharmacokinetics of these drugs were observed. *Effect of Other Drugs on the Metabolism of ARICEFT<sup>®</sup>*: Keloconazole and quinidine, inhibitors of CYP450, 3A4 and 2D6, respectively, inhibit donepazil metabolism *in vitro*. Whether there is a clinical effect of these inhibitors is not known. Inducers of CYP 2D6 and CYP 3A4 (eg, phenytoin, carbamazepine, dexamethasone, rifampin, and phenobarbital) could increase the rate of elimination of ARICEPT\*. Formal pharmacokinetic studies demonstrated that the metabolism of ARICEPT\* is not significantly affected by concurrent administration of digoxin or cimetidine. Use with Anticholinergies: Because of their mechanism of action, cholinesterase inhibitors have the potential to interfere with the activity of anticholinergic their mechanism of use with Cholinesities inhibitors is may be potential to interfere with the activity of anticohinergic medications. Use with Cholinomimetics and Other Cholinesierase inhibitors: A synergistic effect may be expected when cholinergic agonists such as bethanechol. Carcinagenesis, Mutagenesis, Impairment of Fertility Carcinogenicity studies of donepazil have no been completed. Donepezi was not mutagenic in the Ames reverse mutation assay in bacteria. In the chromosome aberration test in cultures of Chinese hamster lung (CHL) cells, some cleaterealio difference and Depezil have need adonteering in the Ames reverse clastogenic effects were observed. Donepezil was not clastogenic in the *in vivo* mouse micronucleus test. Donepezil had no effect on fertility in rats at doses up to 10 mg/kg/day (approximately 8 times the maximum recommended human no effect on fertility in rats at doses up to 10 mg/kg/day (approximately 8 times the maximum recommended human dose on a mg/m<sup>2</sup> basis). **Pregnancy** *Pagnancy Category C:* Teratology studies conducted in pregnant rats at doses up to 16 mg/kg/day (approximately 13 times the maximum recommended human dose on a mg/m<sup>2</sup> basis) and in pregnant rats at the maximum recommended human dose on a mg/m<sup>2</sup> basis) did not disclose any evidence for a tractogenic potential of donepezil. However, in a study in which pregnant rats were given up to 10 mg/kg/day (approximately 16 times the maximum recommended human dose on a mg/m<sup>2</sup> basis) did not disclose any evidence for a tractogenic potential of donepezil. However, in a study in which pregnant rats were given up to 10 mg/kg/day (approximately 8 times the maximum recommended human dose on a mg/m<sup>2</sup> basis) from day 17 of gestation through day 20 postpartum, there was a slight increase in slight decrease in pup survival through day 4 postpartum at this dose; the next lower dose tested was 3 mg/kg/day. There are no adequate or well-controlled studies in pregnant women. ARICEPT<sup>®</sup> should be used during pregnancy only if the potential benefit justifies the potential risk to the fetus. **Nursing Mothers** It is not known whether donepezil is excreted in human breast milk. ARICEPT<sup>®</sup> has no indication for use in nursing mothers. **Pediatric Use** There are no adequate and well-controlled trials to document the safety and efficacy of AIICEPT<sup>®</sup> in any lines socterring in children. **ADVERSE REACTIONS Adverse Events Leading to Discontinuation** The rates of discontinuation from controlled clinical trials of ARICEPT<sup>®</sup> due to adverse events for the ARICEPT<sup>®</sup> 5 mg/day treatment groups were comparable to those of placebo-treatment groups at approximately 5%. The rate for discontinuation of 10 mg/day, was higher at of discontinuation of patients who received 7-day escalations from 5 mg/day to 10 mg/day, was higher at 13%. The most common adverse events leading to discontinuation, defined as those occurring in at least 2% of patients and at twice the incidence seen in placebo patients were nausea (1% [5 mg] and 3% [10 mg] vs 1%

	No ti	tration	One-week titration	Six-week titration	
Adverse Event	Placebo (n=315)	5 mg/day (n=311)	10 mg/day (n=315)	10 mg/day (n=269)	
Nausea	6%	5%	19%	6%	
Diarrhea	5%	8%	15%	9%	
Insomnia	6%	6%	14%	6%	
Fatigue	3%	4%	8%	3%	
Vomiting	3%	3%	8%	5%	
Muscle Cramps	2%	6%	8%	3%	
Anorexia	2%	3%	7%	3%	

[placebo]), diarrhea (<1% [5 mg] and 3% [10 mg] vs 0% [placebo]), and vomiting (<1% [5 mg] and 2% [10 mg] vs <1% [placebo]). Most Frequent Adverse Cilnical Events Seen in Association with the Use of ARICEPT\* The most common adverse events, defined as those occurring at a frequency of at least 5% in patients receiving 10 mg/day and twice the placebo rate, are largely predicted by ARICEPT\*s cholinomimetic effects. These include nausea, diarrhea, insomnia, vomiting, muscle cramp, fatigue, and anoreai. These adverse events were often of mild intensity and transient, resolving during continued ARICEPT\* transment without the need for dose modification. There is evidence to suggest that the frequency of these common adverse events may be affected by the rate of thirston. An open-label study was conducted with 269 patients who received placebo in the 15- and 30-week studies. These patients were filtered to a dose of 10 mg/day over a 6-week period. The rates of common adverse events may be affected by the rate of thrate the comparable to those seen in patients in trade to 10 mg/day over a 6-week period. The rates of common adverse events fraughts and were comparable to those seen in patients on 5 mg/day. See Table 1 for a comparison of the most common adverse events following one week and six week iteriator regimens. Adverse Events Reported in Controlled Trials and were conditions of use, reporting behavior, and the kinds of patients threated may differ. Table 2 lists treatment mergent signs and symptoms that were reported in at least 2% of patients in placebo-controlled trials in the received and the kinds of patients in placebo-controlled trials the received ARICEPT\* and for which the rate of cocurrence was greater for ARICEPT\* assigned than placebo assigned patients. In general, adverse events more hadrese events and were servents of the were terment emergent signs and symptoms that were reported in at least 2% of patients in placebo-controlled trials the received assigned patients.

#### EL220A97

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#### Table 2. Adverse Events Reported in Controlled Clinical Trials in at Least 2% of Patients Receiving ARICEPT\* and at a Higher Frequency They Disable Jested Patients

Body System/Adverse Event	Placebo (n=355)	ARICEPT* (n=747) 74	
Percent of Patients With Any Adverse Event	72		
Body as a Whole			
Headache	9	10	
Pain, Various Locations	8	9	
Accident	6	7	
Fatigue	3	5	
Cardiovascular System		1	
Syncope	1	2	
Digestive System			
Nausea	6	11	
Diarrhea	5	10	
Vomiting	3	5	
Anorexia	2	4	
Hemic and Lymphatic System			
Ecchymosis	3	4	
Metabolic and Nutritional Systems		1	
Weight Decrease	1	3	
Musculoskeletal System			
Muscle Cramps	2	6	
Arthritis	1	2	
Nervous System			
Insomnia	6	9	
Dizziness	6	8	
Depression	<1	3	
Abnormal Dreams	0	3	
Somnolence	<1	2	
Urogenital System			
Frequent Urination	1	2	

Observed During Clinical Trials ARICEPT\* has been administered to over 1700 individuals during clinical trials worldwide. Approximately 1200 of these patients have been treated for at least 3 months and more than 1000 patients have been treated for at least 6 months. Controlled and uncontrolled trials in the United States than 1000 patients have been treated for at least 6 months. Controlled and uncontrolled trials in the United States included approximately 900 patients. In regards to the highest dose of 10 mg/day, this population includes 650 patients treated for 3 months, 475 patients treated for 6 months and 116 patients treated for over 1 year. The range of patient exposure is from 1 to 1214 days. Treatment emergent signs and symptoms that occurred during 3 controlled clinical trials and two open-label trials in the United States were recorded as adverse events by the clinical investigators using terminology of their own choosing. To provide an overal estimate of the proportion of individuals having similar types of events, the events were grouped into a smaller number of standardized categories using a modified COSTART dictionary and event frequencies were calculated across all studies. These categories using a modified to State for the superior and event frequencies were calculated the trials trials where statements are the histories have the treatment on 400 modified to the proportion of the students from these trials were the students of a 000 modified to the studies. These students are used in the listing balaw. The treatment is a studies the studies three to the student to the studies. These trials were an observed to the studies to the studies. These studes the studies the studies three to the studies the studies. These trials were the studies to the studies. These trials were the studies three to the studies the studies the studies. These the students are studies to the studies the studies the studies. 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Events are classified by body system and listed using the following definitions: *trequent* adverse events—those occurring in at least 1/100 patients; *intrequent adverse events*—those occurring in 1/1000 1/1000 patients. These adverse events are not necessarily related to ARICEPT\* treatment and in most cases were events were seen in studies conducted outside the United States. **Body as a Whole**: *Frequent*: influenz, obest patient botherse dreament in the other outside the outside the Bordy as a **Whole**: *Frequent*: influenz, obest patient botherse dreament in the other outside botherse events are not necessarily related to the studies. No important additional adverse events were seen in studies conducted outside the United States. **Body as a Whole**: *Frequent*: influenz, obest patient botherse dreament in the other outside botherse events are not necessarily adverse events budies. No important additional adverse events were seen in studies conducted outside the United States. **Body as a Whole**: *Frequent*: influenze adverse to the studies. No important tadditional adverse events were seen in studies conducted outside the United States. **Body as a Whole**: *Frequent*: influence there are adverse to the studies. No important additional adverse toothache: infrequent: tever, edema face, periorbital edema, hernia hiatal, abscess, cellulitis, chills, generalized coldness, head fullness, listlessness. Cardiovascular System: Frequent: hypertension, vasodilation, atrial Cordiness, field officies, instessifiess. Cardiovascular System: Frequent, Produetti, uppertension, radomation, and hforillation, hot flashes, hypotension, *infrequent*: angina pectoris, postural Mypotension, myocardial infarction, AV block (first degree), congestive heart failure, arteritis, bradycardia, peripheral vascular disease, supraventricular tachycardia, deep vein thrombosis. Dipostive System: Frequent: tecal incontinence, gastrointestinal bleeding, bloating, epigastric pain; *Infrequent*: eructation, gingivitis, increased appelle, flatulence, periodontal abscess, cholettis, increased transaminases, hemortholds, ileus, increased thirst, jaundice, meleand, polydystia, distress, gastroenteritis, increased transaminases, hemortholds, ileus, increased bilita, politis, jaundice, meleand, polydystia, distress, gastroenteritis, increased transaminases, hemortholds, ileus, increased bilita, politis, jaundice, meleand, polydystia, distress, gastroenteritis, increased transaminases, hemortholds, ileus, increased bilita, politis, jaundice, meleand, polydystia, distress, gastroenteritis, processed, transaminases, hemortholds, ileus, increased bilita, politis, jaundice, meleand, polydystia, distress, gastroenteritis, processed, transaminases, hemortholds, ileus, increased bilita, politis, jaundice, meleand, polydystia, distress, gastroenteritis, processed, transaminases, hemortholds, ileus, increased transaminased, hemortholds, distress, gastroenteritis, processed, transaminases, hemortholds, ileus, increased bilita, politis, distress, tasking, polydystia, distress, gastroenteritis, distress, polydystia, distress, gastroenteritis, distress, distres United so that the second s fracture: Infrequent: muscle weakness, muscle tasciculation. Nervoes System: Frequent delusions, tremor, irritability, paresthesia, aggression, vertigo, ataxia, increased libido, restlessness, abnormal crying, nervousness, aphasia; Infrequent: cerebrovascular accident, intracranial hemorrhage, transient ischemic attack, emotional lability, neuralgia, coidness (localized), muscle spasm, dysphoria, gaft abnormality, hypertonia, hypokinesia, neurodermatilits, numbness (localized), paranoia, dysarthria, dysphasia, hostility, decreased libido, melancholia, emotional withdrawal, nystagmus, pacing. Respiratory System: Frequent: dyspnea, sore throat, bronchilts; Infrequent: epistaxis, postnasal drip, pneumonia, hyperventilation, pulmonary congestion, wheezing, hypoxia, pharyngitts, pleurisy, pulmonary collapse, sleep apnea, snoring, Skin and Appendages: Frequent: puritus; daphortesis, uriticaria; Infrequent: dreyens, glaucoma, earche, tinnitus, blepharitis, decreased haring, retinal hemorrhage, otitis externa, otitis media, bad taste, conjunctival hemorrhage, ear buzzing, motion sickness, spots before eyes. Urogenital System: Frequent: urinary incontinence, nocturia; Infrequent dyesus, postagents, urinary urgency, wretorrhagia, cystitis, eurossis, prostate hypertrophy, pederophytik, instriae, neiradenossi, postare hypertrophyterity, bladder, bradenossi, postares, prequent; day and taste, conjunctival hemorrhage, ear buzzing, motion sickness, spots before eyes. Urogenital System: Frequent: urinary incontinence, nocluria; lintrequent: dysuria, hematuria, urinary urgendy, metrorntagia, cystiis, enuresis, prostate hypertophy, pelonephittis, inability to empty bladde, breast fibroadenosis, fibrocystic breast, mastitis, pyuria, renal failure, vaginitis. PostIntroduction Reports Voluntary reports of adverse events temporally associated with ARICEPTP that have been received since market introduction that are not listed above, and that may have no causal relationship with the drug include that followin; a bdominal pain, agitation, cholecystitis, confusion, convulsions, hallucinations, pancreatitis, and rash. OVERDOSAGE Because strategies for the management of overdose are continually evolving, it is advisable to contact a Policon Control Center to determine the latest recommendations of the management of an overdose of any drug. As in any case of overdose, general supportive measures should be utilized. Overdosage with cholinesterias inhibitors can credit in obligancie incide photepriced by eaver a muses uporting saluration, superting hyportaverting photephilos and photephilos. can result in cholinergic crisis characterized by severe nausea, vomiting, salivation, sweating, bradycardia, hypotension, respiratory depression, collapse and convulsions. Increasing muscle weakness is a possibility and may result in death if respiratory muscles are involved. Tertiary anticholinergics such as atropine may be used as an antidote for ARICEPT® overdosage. Intravenous atropine sulfate titrated to effect is recommended: an initial dose of 1.0 antidote for ARICEPT\* overdosage. Intravenous atropine sulfate titrated to effect is recommended: an initial dose of 1.0 to 2.0 mg IV with subsequent doses based upon clinical response. Alypical responses in blood pressure and heart rate have been reported with other cholinomingtics when co-administered with quaternary anticholinergics such as glycopyrrolate. It is not known whether ARICEPT\* and/or its metabolites can be removed by dialysis (hemodialysis, peritoneal dialysis, or hemofiltration). Dose-related signs of toxicity in animals included reduced spontaneous movement, prone position, staggering gait, lacimation, clonic convulsions, depressed respiration, salivation, miosis, termors, fasciculation and lower body surface temperature. **DOSAGE AND ADMINISTRATION** The dosages of ARICEPT\* shown to be effective in controlled clinical triats are 5 mg and 10 mg administered once per day. Controlled chicate hat the 10 mg dose, with a one week titration, is likely to be associated with a higher incidence of contemplated until patients have been on a daily dose of 15 mg for 4 to 6 weeks. Whether or not to employ a dose of 10 mg is a matter of prescriber and patient preference. ARICEPT\* should be taken in the evening, just prior to retiring, and may be taken with or without food.



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# **CNS SPECTRUMS**

## The International Journal of Neuropsychiatric Medicine

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## IN THE JOURNAL OF MARCH 1998

"Autism is as common as many other childhood medical disorders such as diabetes and leukemia. However, detection of the autistic child is often delayed significantly beyond the time of first suspicion by parents."

## A DIMENSIONAL PERSPECTIVE ON AUTISM

#### page 23

"The task of linking symptomatology and behavioral patterns with neurobiological abnormalities in a genetically influenced developmental disorder is formidable. The integration of research findings is necessary when attempting to find convergence of evidence linking clinical, functional, and neuroanatomical abnormalities in autism. By using a dimensional perspective to identify and clarify the specific components that give rise to the autistic syndrome, we are in a better position to determine the neurobiological determinants of the specific core autistic dimensions. This approach attempts to solve the problems of etiologic heterogeneity, developmental variation, and neurological comorbidity."

## <u>GUIDELINES FOR EARLY DIAGNOSIS</u> page 48

"Autism is as common as many other childhood medical disorders such as diabetes and leukemia. However, detection of the autistic child is often delayed significantly beyond the time of first suspicion by parents. The CHAT is a rapid and convenient tool that should be used to screen all 18-month-old children for autistic spectrum disorders....This consensus statement highlights the large gaps in our current knowledge regarding the appropriate evaluation of children with autism and related disorders (PDD). At this point, management depends largely upon the expert judgment of clinicians experienced in autism, language delay, and pediatric neurology and psychiatry, rather than data gathered from controlled clinical trials."

## NEUROPATHOLOGICAL <u>FINDINGS IN AUTISM</u> page 54

"The report of smallness of the brainstem in some studies, but not in others, suggests brainstem involvement in autism. Depletion of cells in the facial nucleus and abnormalities of the hypoglossal nucleus and superior olive were described in the brainstem of a 21-year-old woman with autism in whom the complement of neurons in the neocortex was normal. It is unfortunate that a detailed neurologic examination was not available, since an occasional individual with Möbius' syndrome, characterized by dysgenesis of multiple cranial nerve nuclei, most often the VIth and VIIth, has autistic behaviors. As most children with Möbius' syndrome are not autistic, the relevance of these findings in the cranial nerve nuclei remains unclear."

#### THE AUTISM AND EPILEPSY <u>CONNECTION</u> page 61

"The variability in seizure frequency reported in ASD is thus likely due to three factors: (1) the age groups studied, with the highest percent of seizures found in studies that included adolescents and young adults; (2) the level of cognitive function, with the highest percent of seizures found in studies that included children with severe mental deficiency; and (3) the type and degree of language dysfunction, with the highest percent of seizures occurring in individuals with VAA."

## THE IMMUNOLOGIC THEORY page 71

"Autism is a syndrome resulting from several different etiologies or a combination of pathological mechanisms. Recent studies, to be reviewed later, suggest that immune dysfunction may be one contributing factor to the development of some cases of this severe developmental disorder. According to this theory, some children may be susceptible to an environmental pathogen (most likely a virus or bacterium) resulting from an inherited deficiency of their immune system. This deficiency could result from decreased T-cell-mediated immunity, decreased levels of an immunoglobulin (Ig) such as IgA, or a nonspecific defense system including complement activity. Unable to clear the pathogen in a timely and normal manner, the child is at increased risk for the pathogen (or pathogenic toxins) to damage the developing brain and cause the symptoms of autism."

#### THE CEREBELLUM'S INFLUENCE ON <u>MENTAL IMAGERY</u> page 80

"Besides neurochemistry, neuroanatomical studies involving autopsied brain samples from autistic patients have produced evidence of alterations in the hippocampus, amygdala, and cerebellum; in particular at this level, a loss in Purkinje's cells has been detected....The cerebellum has been considered for decades as contributing only to motor coordination and control. In recent years, however, the theory has emerged that it may have a role in cognition, emotional processes, and internal mental imagery."

## SCHOOL'S IN PROGRESS. HOW ARE YOUR PATIENTS PROGRESSING ON THE ADHD TREATMENT REGIMENS YOU PRESCRIBED?

## PRESCRIBE ADDERALL<sup>®</sup>-IT MAY MAKE A DIFFERENCE

As children settle into the routine of a structured classroom environment and teachers become more familiar with individual capabilities and behavior patterns, potential problem behavior and academic underachievement may become more apparent. A change in medication may be warranted to optimize individual ADHD treatment plans.

## The ADDERALL® (mixed salts of a single-entity amphetamine product) Formulation and Starting Dosage Frequency of One to Two Times Per Day' May Make a Difference

**ADDERALL** is the only ADHD product available to contain both dextro (*d*) and levo (*l*) amphetamine **ADDERALL** usage data (n=611) indicate that **OVER 90% OF PATIENTS** can be maintained on a dosage frequency of one to two times per day<sup>2\*</sup>

**ADDERALL** usage data (n=611) indicate that most patients, across a range of doses, do not experience adverse events with a frequency of more than  $1\%^{2*}$ 

**ADDERALL** is available in 5 mg, 10 mg, 20 mg, and **NEW 30 mg** double-scored tablets which allows you to achieve precise dosage correlation with individual therapeutic needs in a single prescription

As with most psychostimulants indicated for ADHD, the possibility of growth suppression and the potential for precipitating motor tics and Tourette's syndrome exists with ADDERALL treatment, and in rare cases exacerbations of psychosis have been reported. Since amphetamines have a high potential for abuse, ADDERALL should only be prescribed as part of an overall multimodal treatment program for ADHD with close physician supervision.



\*Thirty-four patients receiving greater than 40 mg per day were excluded from this analysis.

Please see reverse side for references and brief summary of prescribing information.

REFERENCES: I. ADDERALL Package Insert, Richwood Pharmaceutical Company Inc. 2. Data on file, Richwood Pharmaceutical Company Inc. Analysis of open-label data collected from March 1995 through February 1996.



## ADDERALL<sup>®</sup> TABLETS

AMPHETAMINES HAVE A HIGH POTENTIAL FOR ABUSE. ADMINISTRATION OF AMPHETAMINES FOR PROLONGED PERIODS OF TIME MAY LEAD TO DRUG DEPENDENCE AND MUST BE AVOIDED. 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.

INDICATIONS: Attention Deficit Disorder with Hyperactivity: ADDERALL is indicated as an integral part of a total treatment program which typically includes other remedial measures (psychological, educational, social) for a stabilizing effect in children with behavioral syndrome characterized by the following group of developmentally inappropriate symptoms: moderate to severe distractibility, short attention span, hyperactivity, emotional lability, and impulsivity. The diagnosis of this syndrome should not be made with finality when these symptoms are only of comparatively recent origin. Nonlocalizing (soft) neurological signs, learning disability and abnormal EEG may or may not be present, and a diagnosis of central nervous system dysfunction may or may not be warranted. In Narcolepsy: CONTRAINDICATIONS: Advanced arteriosclerosis, symptomatic cardiovascular disease, moderate to severe hypertension, hyperthyroidism, known hypersensitivity or idiosyncrasy to the sympathomimetic 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: Clinical experience suggests that in psychotic children, administration of amphetamine may exacerbate symptoms of behavior disturbance and thought disorder. Data are inadequate to determine whether chronic administration of amphetamine may be associated with growth inhibition; therefore, growth should be monitored during treatment. Usage in Nursing Mothers: Amphetamines are excreted in human milk. Mothers taking amphetamines should be advised to refrain from nursing. PRECAUTIONS: General: Caution is to be exercised in prescribing amphetamines for patients with even mild hypertension. The least amount feasible should be prescribed or dispensed at one time in order to minimize the possibility of overdosage. Information for Patients: Amphetamines may impair the ability of the patient to engage in potentially hazardous activities such as operating machinery or vehicles; the patient should therefore be cautioned accordingly. Drug Interactions: Acidifying agents -Gastrointestinal acidifying agents (guanethidine, reserpine, glutamic acid HCI, ascorbic acid, fruit juices, etc.) lower absorption of amphetamines. Urinary acidifying agents -(ammonium chloride, sodium acid phosphate, etc.) Increase the concentration of the ionized species of the amphetamine molecule, thereby increasing urinary excretion. Both groups of agents lower blood levels and efficacy of amphetamines. Adrenergic blockers - Adrenergic blockers are inhibited by amphetamines. Alkalinizing agents - Gastrointestinal alkalinizing agents (sodium bicarbonate, etc.) increase absorption of amphetamines. Urinary alkalinizing agents (acetazolamide, some thiazides) increase the concentration of the nonionized species of the amphetamine molecule, thereby decreasing urinary excretion. Both groups of agents increase blood levels and therefore potentiate the actions of amphetamines. Antidepressants, tricyclic - Amphetamines may enhance the activity of tricyclic or sympathomimetic agents; d-amphetamine with desipramine or protriptyline and possibly other tricyclics cause striking and sustained increases in the concentration of d-amphetamine in the brain; cardiovascular effects can be potentiated. MAO inhibitors - MAOI antidepressants, as well as a metabolite of furazolidone, slow amphetamine metabolism. This slowing potentiates amphetamines, increasing their effect on the release of norepinephrine and other monoamines from adrenergic nerve endings; this can cause headaches and other signs of hypertensive crisis. A variety of neurological toxic effects and malignant hyperpyrexia can occur, sometimes with fatal results. Antihistamines -Amphetamines may counteract the sedative effect of antihistamines. Antihypertensives Amphetamines may antagonize the hypotensive effects of antihypertensives. Chlorpromazine - Chlorpromazine blocks dopamine and norepinephrine reuptake, thus inhibiting the central stimulant effects of amphetamines, and can be used to treat amphetamine poisoning. Ethosuximide - Amphetamines may delay intestinal absorption of ethosuximide. Haloperidol - Haloperidol blocks dopamine and norepinephrine reuptake, thus inhibiting the central stimulant effects of amphetamines. Lithium carbonate - The anorectic and stimulatory effects of amphetamines may be inhibited by lithium carbonate. Meperidine Amphetamines potentiate the analgesic effect of meperidine. Methenamine therapy -Urinary excretion of amphetamines is increased, and efficacy is reduced, by acidifying agents used in methenamine therapy. Norepinephrine - Amphetamines enhance the adrenergic effect of norepinephrine. Phenobarbital - Amphetamines may delay intestinal absorption of phenobarbital; co-administration of phenobarbital may produce a synergistic anticonvulsant action. Phenytoin - Amphetamines may delay intestinal absorption of phenytoin; co-administration of phenytoin may produce a synergistic anticonvulsant action. Propoxyphene - In cases of propoxyphene overdosage, amphetamine 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: Mutagenicity studies and long-term studies in animals to determine the carcinogenic potential of amphetamine, have not been performed. **Pregnancy - Teratogenic Effects:** Pregnancy Category C. Amphetamine has been shown to have embryotoxic and teratogenic effects when administered to A/Jax mice and C57BL mice in doses approximately 41 times the maximum human dose. Embryotoxic effects were not seen in New Zealand white rabbits given the drug in doses 7 times the human dose nor in rats given 12.5 times the maximum human dose. While there are no



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adequate and well-controlled studies in pregnant women, there has been one report of severe congenital bony deformity, tracheoesophageal 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 risk to the fetus. Nonteratogenic Effects: Infants born to mothers dependent on amphetamines have an increased risk of premature delivery and low birth weight. Also, these infants may experience symptoms of withdrawal as demonstrated by dysphoria, including agitation, and significant lassitude. Pediatric Use: Long-term effects of amphetamines in children have not been well established. Amphetamines are not recommended for use in children under 3 years of age with Attention Deficit Disorder with Hyperactivity described under INDICATIONS AND USAGE. 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. Drug treatment is not indicated in all cases of Attention Deficit Disorder with Hyperactivity and should be considered only in light of the complete history and evaluation of the child. The decision to prescribe amphetamines should depend on the physician's assessment of the chronicity and severity of the child's symptoms and their appropriateness for his/her age. Prescription should not depend solely on the presence of one or more of the behavioral characteristics. When these symptoms are associated with acute stress reactions, treatment with amphetamines is usually not indicated. ADVERSE REACTIONS: Cardiovascular: Palpitations, tachycardia, elevation of blood pressure. There have been isolated reports of cardiomyopathy associated with chronic amphetamine use. Central Nervous System: Psychotic episodes at recommended doses (rare), overstimulation, restlessness, dizziness, insomnia, euphoria, dyskinesia, dysphoria, tremor, headache, exacerbation of motor and phonic tics and Tourette's syndrome. Gastrointestinal: Dryness of the mouth, unpleasant taste, diarrhea, constipation, other gastrointestinal disturbances. Anorexia and weight loss may occur as undesirable effects when amphetamines are used for other than the anorectic effect. Allergic: Urticaria. Endocrine: Impotence, changes in libido. DRUG ABUSE AND DEPENDENCE: Dextroamphetamine sulfate 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 include severe dermatoses, marked insomnia, irritability, hyperactivity, and personality changes. The most severe manifestation of chronic intoxication is psychosis, often clinically indistinguishable from schizophrenia. This is rare with oral amphetamines. OVERDOSAGE: Individual patient response to amphetamines varies widely. While toxic symptoms occasionally occur as an idiosyncrasy at doses as low as 2 mg, they are rare with doses of less than 15 mg; 30 mg can produce severe reactions, yet doses of 400 to 500 mg are not necessarily fatal. In rats, the oral LD50 of dextroamphetamines ulfate is 96.8 mg/kg. Symptoms: Manifestations of acute overdosage with amphetamines include restlessness, tremor, hyperreflexia, rapid respiration, confusion, assaultiveness, hallucinations, panic states, hyperpyrexia and rhabdomolysis. Fatigue and depression usually follow the central stimulation. Cardiovascular effects include arrhythmias, hypertension or hypotension and circulatory collapse. Gastrointestinal symptoms include nausea, vomiting, diarrhea, and abdominal cramps. Fatal poisoning is usually preceded by convulsions and coma. Treatment: Consult with a Certified Poison Control Center for up to date guidance and advice. Management of acute amphetamine intoxication is largely symptomatic and includes gastric lavage, administration of activated charcoal, administration of a cathartic and sedation. Experience with hemodialysis or peritoneal dialysis is inadequate to permit recommendation in this regard. Acidification of the urine increases amphetamine excretion, but is believed to increase risk of acute renal failure if myoglobinuria is present. If acute, severe hypertension complicates amphetamine overdosage, administration of intravenous phentolamine (Regitine\*, CIBA) has been suggested. However, a gradual drop in blood pressure will usually result when sufficient sedation has been achieved. Chlorpromazine antagonizes the central stimulant effects of amphetamines and can be used to treat amphetamine intoxication. DOSAGE AND ADMINISTRATION: Regardless of indication, amphetamines should be administered at the lowest effective dosage and dosage should be individually adjusted. Late evening doses should be avoided because of the resulting insomnia. Attention Deficit Disorder with Hyperactivity: Not recommended for children under 3 years of age. In children from 3 to 5 years of age, start with 2.5 mg daily; daily dosage may be raised in increments of 2.5 mg at weekly intervals until optimal response is obtained. In children 6 years of age and older, start with 5 mg once or twice daily; daily dosage may be raised in increments of 5 mg at weekly intervals until optimal response is obtained. Only in rare cases will it be necessary to exceed a total of 40 mg per day. Give first dose on awakening; additional doses (1 or 2) at intervals of 4 to 6 hours. Where possible, drug administration should be interrupted occasionally to determine if there is a recurrence of behavioral symptoms sufficient to require continued therapy. Narcolepsy: Usual dose 5 mg to 60 mg per day in divided doses, depending on the individual patient response. Narcolepsy seldom occurs in children under 12 years of age; however, when it does dextroamphetamine sulfate, may be used. The suggested initial dose for patients aged 6-12 is 5 mg daily; daily dose may be raised in increments of 5 mg at weekly intervals until optimal response is obtained. In patients 12 years of age and older, start with 10 mg daily; daily dosage may be raised in increments of 10 mg at weekly intervals until optimal response is obtained. If bothersome adverse reactions appear ( e.g., insomnia or anorexia), dosage should be reduced. Give first dose on awakening; additional doses (1 or 2) at intervals of 4 to 6 hours. CAUTION: Federal law prohibits dispensing without prescription.



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**References:** American Medical Association style. See the following examples:

- 1. Jones J. Necrotizing Candida esophagitis. JAMA. 1980;244:2190-2191.
- 2. Stryer L. *Biochemistry*. 2nd ed. San Francisco, Calif: WH Freeman Co; 1980:559-596.

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PAXIL® (brand of paroxetine hydrochloride) See complete prescribing information in SmithKline Beecham Pharmaceuticals literature or PDR. The following is a brief summary. INDICATIONS AND USAGE: Paxil is indicated for the treatment of depression, obsessions and com-

INDICATIONS AND USAGE: Paxil is indicated for the treatment of depression, obsessions and compulsions in patients with obsessive compulsive disorder (OCD) as defined in DSM-IV, and panic disorder, with or without agoraphobia, as defined in DSM-IV.
 CONTRAINDICATIONS: Concomitant use in patients taking monoamine oxidase inhibitors (MAOIs) is contraindicated. (See WARNINGS and PRECAUTIONS.)
 WARNINGS: Interactions with MAOIs may occur. Given the fatal interactions reported with concomitant or immediately consecutive administration of MAOIs and other SSRIs, do not use Paxil in combination with a MAOI or within 2 weeks of discontinuing MAOI treatment. Allow at least 2 weeks after stopping Paxil before starting a MAOI.
 PRECAUTIONS: As with all antidepressants, use Paxil cautiously in patients with a history of mania. Use Paxil cautiously in patients with a history of seizures. Discontinue it in any patient who develops seizures.

Sources. The possibility of suicide attempt is inherent in depression and may persist until significant remission occurs. Close supervision of high-risk patients should accompany initial drug therapy. Write *Paxil* pre-scriptions for the smallest quantity of tablets consistent with good patient management in order to reduce the risk of overdose.

Reversible hyponatremia has been reported, mainly in elderly patients, patients taking diuretics or those who were otherwise volume depleted. Abnormal bleeding (mostly ecchymosis and purpura), including a case of impaired platelet aggregation, has been reported; the relationship to paroxetine is unclear. Clinical experience with *Paxil* in patients with concomitant systemic illness is limited. Use cautiously in

patients with diseases or conditions that could affect metabolism or hemodynamic responses. Observe the usual cautions in cardiac patients. In patients with severe renal impairment (creatinine clearance

the usual cautions in cardiac patients. In patients with severe renal impairment (creatinine clearance <30 mL/min.) or severe hepatic impairment, a lower starting dose (10 mg) should be used. Caution patients about operating hazardous machinery, including automobiles, until they are reasonably sure that *Paxil* therapy does not affect their ability to engage in such activities. Tell patients 1) to con-tinue therapy as directed; 2) to inform physicians about other medications they are taking or plan to take; 3) to avoid alcohol while taking *Paxil*; 4) to notify their physicians if they become pregnant or intend to become pregnant during therapy, or if they're nursing. Weakness, hyperreflexia, and incoordination following use of an SSRI and sumatriptan have been rarely

reported.

reported. Concomitant use of *Paxil* with tryptophan is not recommended. Use cautiously with warfarin. When ad-ministering *Paxil* with cimetidine, dosage adjustment of *Paxil* after the 20 mg starting dose should be guided by clinical effect. When co-administering *Paxil* with phenobarbital or phenytoin, no initial *Paxil* dosage adjustment is needed; base subsequent-changes on clinical effect. Concomitant use of *Paxil* with drugs metabolized by cytochrome  $P_{adj}ID_{i}$  (antidepressants such as nortriptyline, antirptyline, impramine, designamine and fluoxetine; phenothiazines such as thioridazine; Type 10 antirptyline) impramine, designamine and fluoxetine; phenothiazines such as thioridazine; type IC antarrhythmics such as propafenone, fecainide and encainide) or with drugs that inhibit this enzyme (e.g., quinidine) may require lower doses than usually prescribed for either *Paxil* or the other drug; approach concomi-tant use cautiously. An *in vivo* interaction study revealed that paroxetine had no effect on terfenadine pharmacokinetics. Additional *in vitro* studies showed that the inhibitory effects of paroxetine on other IIIA<sub>s</sub> substrates (astemizole, cisapride, triazolam and cyclosporin) was at least 100 times less potent than ketoconazole, a potent IIIA<sub>s</sub> inhibitor. Assuming that the relationship between paroxetine's *in vitro* Ki and its lack of effect on terfenadine's *in vivo* clearance predicts its effect on other IIIA<sub>s</sub> substrates. Ki and its lack of effect on terfenadines *in vivo* clearance predicts its effect on other IIIA, substrates, paroxetine's inhibition of IIIA, activity should have little clinical significance. Use caution when co-administering *Paxil* with tricyclic antidepressants (TCAs). TCA plasma concentrations may need moni-toring and the TCA dose may need to be reduced. Administration of *Paxil* with another tightly protein-bound drug may shift plasma concentrations, resulting in adverse effects from either drug. Concomitant use of *Paxil* and alcohol in depressed patients is not advised. Undertake concomitant use of *Paxil* and lithium or digoxin cautiously. If adverse effects are seen when co-administering *Paxil* with procyclidine, reduce the procyclidine dose. Elevated theophylline levels have been reported with *Paxil* co-administra-tion, monitoring theoremilium levels.

In a constraint of the provide the constraint of the constraint of the constraint of the provide the provide the constraint of the constra evidence of mutagenicity with Paxil.

Rats receiving paroxetine at 15 mg/kg/day (2.4 times the MRHD on a mg/m<sup>2</sup> basis) showed a reduced pregnancy rate.

hats feedbilling partocentile at 15 mg/kg/04y/2.4 times the MnHb off a ling/in "basis) showed a reduced preg-pancy rate. **Pregnancy Category C.** Reproduction studies performed in rats and rabbits at doses up to 6 mg/kg/day, 8.1 (rat) and 1.9 (rabbit) times the MRHD on a mg/m<sup>2</sup> basis, have revealed no evidence of teratogenic effects or of selective toxicity to the fetus. However, rat pup deaths increased during the first 4 days of lactation when dosing occurred during the last trimester of gestation and continued throughout lac-tation. The cause of these deaths is not known. There are no adequate and well-controlled studies in pregnant women. *Paxil* should be used in pregnancy only if the potential benefit justifies the potential risk to the fetus. The effect of *Paxil* on labor and delivery in humans is unknown. Paroxetine is secreted in human milk, exercise caution when administering *Paxil* to a nursing woman. Safety and effectiveness in the pediatric population have not been established. In worldwide premarketing *Paxil* clinical trials, 17% of *Paxil*-treated patients were ≥65 years of age. Pharmacokinetic studies revealed a decreased clearance in the elderyt, however, there were no overall differences in the adverse event profile between older and younger patients. **ADVERSE REACTIONS: Incidence in Controlled Trials—Commonly Observed Adverse Events in Controlled Clinical Trials:** The most commonly observed adverse events associated with the use of *Paxil* in the treatment of depression (incidence of 5% or greater and incidence for *Paxil* at least twice that for placebol; asthenia (15% vs. 6%), sweating (11% vs. 2%), insomnia (13% vs. 6%), decreased appetite (6% vs. 2%), somnolence (23% vs. 9%), dizziness (13% vs. 6%) insomnia (13% vs. 6%), terotomeser (10% vs. 2%).

orders (10% vs. 0%).

orders (10% vs. 0%). The most commonly observed adverse events associated with the use of paroxetine in the treatment of obsessive compulsive disorder (incidence of 5% or greater and incidence for *Paxil* at least twice that of placebo) were: nausea (23% vs. 10%), dry mouth (18% vs. 9%), decreased appetite (9% vs. 3%), constipation (16% vs. 6%), diziness (12% vs. 6%), somolence (24% vs. 7%), tremor (11% vs. 1%), sweating (19% vs. 9%), impotence (8% vs. 19%) and ahormal ejaculation (123% vs. 1%). The most commonly observed adverse events associated with the use of paroxetine in the treatment of paric disorder (incidence of 5% or greater and incidence for *Paxil* at least twice that for placebo) were: asthenia (14% vs. 5%), sweating (14% vs. 6%), decreased appetite (7% vs. 3%), libido decreased (19% vs. 1%), termor (19% vs. 1%), abnormal ejaculation (21% vs. 1%), by, by useding (14% vs. 5%), sweating (14% vs. 6%), decreased appetite (7% vs. 3%), libido decreased (19% vs. 1%), internor (19% vs. 1%), abnormal ejaculation (21% vs. 1%), female genital disorders (9% vs. 1%) and impotence (5% vs. 0%). Twenty percent (1, 199;6, 145) of *Paxil* patients in worldwide clinical trials in depression and 11.8% (64/542) and 9.4% (44/469) of *Paxil* patients in worldwide trials in 0CD and panic disorder, respectively, discontinued treatment to an adverse event. The most common events (>1%) associated with discontinued to an adverse veent chiclude the following: **depression** somnolence, agita-

continuation and considered to be drug related include the following: **depression**-somnolence, agita-tion, tremor, nausea, diarrhea, dry mouth, vomiting, asthenia, abnormal ejaculation, sweating;

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OCD-insomnia, dizziness, constipation, nausea, asthenia, abnormal ejaculation, impotence; panic dis order-somnolence, insomnia, nausea

**Order**-sommolence, insomnia, nausea. The following adverse events occurred in 6-week placebo-controlled trials of similar deadache, asthe-nia, palpitation; vasodilation; sweating, rash; nausea, dry mouth, constipation, diarrhea, decreased appetite, flatulence, oropharynx disorder, dyspepsia; myopathy, myalgia, myasthenia; somolence, dizi-ness, insomnia, tremor, nervousness, anxiety, paresthesia, libido decreased, drugged feeling, confusion; yawn; blurred vision, taste perversion; ejaculatory disturbance, other male genital disorders, urinary fre-quency, urination disorder, female genital disorders. The following adverse events occurred at a frequency of 2% or more among OCD patients on *Paxil* who participated in placebo-controlled trials of 12-weeks duration in which patients were dosed in a range of 20 to 60 mg/day or among patients with panic disorder on *Paxil* who participated in placebo-controlled trials of 12-weeks duration in which patients were dosed in a range of 10 to 12 weeks duration in which patients were dosed in a range of 10 to 12 weeks duration in which patients were dosed in a range of 10 to 12 weeks duration, in disrhea, decreased, agpetite, increased appetite; insomnia, sommolence, dizziness; themor, nervousness<sup>\*\*</sup>, libido decreased, agpitation<sup>\*\*</sup>, seveating, rash<sup>\*\*</sup>, nausea, dry mouth, constipation, diarhea, decreased, agpitation<sup>\*\*</sup>, andrey; ahonrmal dreams<sup>\*\*</sup>, concentration impaired<sup>\*\*</sup>, depersonalization<sup>\*\*</sup>, myoclonus, amnesia<sup>\*\*</sup>, rhinitis<sup>\*</sup>, ahonrmal vision<sup>\*\*</sup>, taste perversion<sup>\*\*</sup>; ahonrmal ejaculation, female genital disorder, impotence, urinary frequency patients end. \*\* denotes occurred at a frequency of patients were dosed in a range of 10 to 50 mg/day or among patients with patients were dosed in a range of 10 to 50 mg/day or among patients with patients were dosed in a range of 10 to 50 mg/day. \*\* ansee, diventions, themory, how setting the patients were dosed in a range of 10 to 50 mg/day or among patients were dosed in a range of 10 to 50 mg/day or a

patients only. Studies show a clear dose dependency for some of the more common adverse events associated with Paxil use. There was evidence of adaptation to some adverse events with continued Paxil therapy (e.g., nausea and dizziness). Significant weight loss may be an undesirable result of Paxil treatment for some patients but, on average, patients in controlled trials had minimal (about 1 lb) loss. In placebo-controlled clinical trials, Paxil-treated patients exhibited abnormal values on liver function tests no more frequentthan placebo-treated patie

Cinical trials, *Paxit* related patients exhibited abnormal values on liver function tests no more frequent-ly than placebo-treated patients. **Other Events Observed During the Premarketing Evaluation of Paxif**: During premarketing as-sessment in depression multiple doses of *Paxii* were administered to 6,145 patients in phase 2 and 3 studies. During premarketing clinical trials in OCD and painc disorder, 542 and 459 patients, respective-ly, received multiple doses of *Paxii*. The following adverse events were reported. Note: "frequent" = events occurring in at least 1/100 patients; "infrequent" = 1/100 to 1/1000 patients; "rare" = less than 1/1000 patients. Events are classified within body system categories and enumerated in order of decreasing frequency using the above definitions. It is important to emphasize that although the events occurred during *Paxii* treatment, they were not necessarily caused by it. **Body as a Whole:** frequent: chills, malaise; infrequent: Hergic reaction, carcinoma, face edema, moniliasis, neck pain; rare: abscess, adrenergic syndrome, cellulitis, neck rigidity, pelvic pain, peritoni-tis, shock, ulcer. **Cardiovaccular System:** *Trequent*: hypertension, syncope, tachycardia; *infrequent*: peripheral vascular disorder; *rare*: angina pectoris, arrhythmia, atrial fibrillation, bundle branch block, cerebral ischemia, cerebrovascular accident, congestive heart failure, heart block, low cardiac output, mycoardial infarct, mycoardial ischemia, pallor, phlebitis, pulmonary embolus, supraventricular extrasys-toles, thrombophlebitis, thrombosis, varicose vein, vascular headache, ventricular extrasys-tists, increased salivation. Iliver function tests abnormal, montulecratin merchale merchage. **Digestive System:** *infrequent*; bruxism, collits, dysphagia, eructation, gastroentertiis, duodenitis, entertiis, bloody diarrhea, bulimia, choleithiasis, duodenitis, entertiis, stris, increased salivation, liver function tests abnormal, mouth ulceration, rectal hemorrhage, ulcerative stomatitis; *rare*: aphthous stomatitis, bloody diarrhea, bulimia, cholelithiasis, duodenitis, enteritis, esophagitis, fecal impactions, fecal incontinence, gastritis, gum hemorrhage, hematemesis, hepatitis, ileus, intestinal obstruction, jaundice, melena, peptic ulcer, salivary gland enlargement, stomach ulcer, stomatitis, tongue discoloration, tongue edema, tooth caries, tooth malformation. Endocrine Systems: *infrequent*: anemia, leukopenia, lymphadenopathy, purpura; *rare*: abnormal erythrocytes, basophilia, eosinophilia, hypochromic anemia, iron deficiency anemia, leukocytosis, lymphedema, abnormal lymphordes, microphile, anemia, thrombocytes; mercortie anemia, thrombocytes; mercorties, anemia, thrombocytes; mercortie anemia, thrombocytes; basophilia, eosinophilia, hypochromic anemia, iron deficiency anemia, leukocytosis, lymphedema, abnormal lym-phocytes, lymphocytosis, microcytic anemia, monocytosis, normocytic anemia, thrombocythemia. Metabolic and Nutritional: frequent: edema, weight gain, weight losis; infrequent: hyperglycemia, peripheral edema, SGOT increased, SGPT increased, thirst; rare: alkaline phosphatase increased, biliru-binemia, BUN increased, creatinine phosphokinase increased, dehydration, gamma globulins increased, gout, hypercalcemia, hyporcholesteremia, hyperkalemia, hyperphosphatemia, hypocalcemia, hypo-glycemia, hypokalemia, hyponatremia, ketosis, lactic dehydrogenase increased. Musculoskeletal System: frequent: arthralgia; infrequent: arthritis; rare: arthrosis, bursitis, myositis, osteoporosis, gen-eralized spasm, tenosynovitis, tetany. Nervous: System: frequent: abnormal thinking, akinesia, alcohol abuse, ataxia, convulsion, depersonalization, dystonia, hallucinations, hostitiliv, hyperkinesia, hypertonia, hypesthesia, incoordination, lack of emotion, manic reaction, neurosis, paralysis, paranoid reaction; rare: abnormal electroencephalogram, abnormal gait, antisocial reaction, aphasia, choreoa-thetosis, circumoral paresthesia, delirium, delusions, diplopia, drug dependence, dysarthira, dyskinesia, hysteria, libido increased, manic-depressive reaction, meningits, myeralgesia, hypokinesia euphoria, extrapyramidal syndrome, fasciculations, grand mal convulsion, hyperaigasia, hypokinesia, hysteria, libido increased, manic-depressive reaction, meningitis, myelitis, neuralgia, neuropathy, nys-tagmus, peripheral neuritis, psychosis, psychotic depression, reflexes decreased, reflexes increased, stupor, trismus, withdrawal syndrome. **Respiratory System:** *traquent*: cough increased, thinitis; *infre-quent*: asthma, bronchitis, dyspnea, epistaxis, hyperventilation, pneumonia, respiratory flu, sinusitis, voice alteration; *rare*: emphysema, hemoptysis, hiccups, lung fibrosis, pulmonary edema, sputum increased. **Skin and Appendages:** *traquent*: pruritus; *infrequent*: acne, alopacia, dry skin, acchrymosis, eczema, furunculosis, uriticaria; *rare*: angloedema, contact dematitis, enthema nodosum, erythema multiforme, fungal dermatitis, herpes simplex, herpes zoster, hirsutism, maculopapular tash, photosen-sitivity, seborrhea, skin discoloration, skin hypertrophy, skin melonoma, skin ulcer, vesiculobullous rash. **Special Senses:** *traquent*: tinnitus; *infrequent*: abnormality of accommodation, conjunctivitis, ear pain, eya pain, mydriasis, otitis media, taste loss, visual field defect; *rare*: amblyopia, anisocoria, blephartits, cataract, conjunctival edema, corneal ulcer, defices, exophthalmos, eve hemorrhage, olaucoma, hypereve pain, mydriasis, otitis media, taste loss, visual field defect; rare: amblyopia, anisocoria, blepharitis, cataract, conjunctival edema, corneal ulcer, deafness, exophthalmos, eye hemorrhage, glaucoma, hyper-acusis, keratoconjunctivitis, night blindness, otitis externa, parosmia, photophobia, ptosis, retinal hem-orrhage. **Urogenital System:** infrequent: abortion, amenorrhea, breast pain, cystitis, dysmenorrhea, dysuria, hematuria, menorrhagia, nocturia, polyuria, urethritis, urinary incontinence, urinary retention, urinary urgency, vaginitis: *rare:* breast atrophy. breast carcinoma, breast enlargement, breast neoplasm, epididymitis, female lactation, fibrocystic breast, kidney calculus, kidney function abnormal, kidney pain, leukorrhea, mastitis, metrorrhagia, nephritis, oliguria, prostatic carcinoma, pyuria, urethritis, uterine spasm, urolith, vaginal hemorrhage, vaginal moniliasis. **Postmarketing Reports** 

Voluntary reports of adverse events that have been received since market introduction and not listed above that may have no causal relationship with *Paxil* include-acute pancreatitis, elevated liver func-tion tests (the most severe cases were deaths due to liver necrosis, and grossly elevated transminas-es associated with severe liver dystuction), Guillain-Barré syndrome, toxic epidemal necrolysis, pri-apism, thrombocytopenia, syndrome of inappropriate ADH secretion, symptoms suggestive of proapism, thrombocytopenia, syndrome of inappropriate ADH secretion, symptoms suggestive of pro-lactinemia and galactorrhea, neuroleptic malignant syndrome-like events, extrapyramidal symptoms which have included akathisia, bradykinesia, cogwheel rigidity, dystonia, hypertonia, oculogyric crisis (which has been associated with concomitant use of pimozide), tremor and trismus; and serotonin syn-drome, associated in some cases with concomitant use of serotonergic drugs and with drugs which may have impaired *Paxil* metabolism (symptoms have included agitation, confusion, diaphoresis, hallucina-tions, hyperreflexia, myoclorus, shivering, tachycardia and tremor). There have been spontaneous re-ports that abrupt discontinuation may lead to symptoms such as dizziness, sensory disturbances, agita-tion or anxiety, nausea and sweating; these events are generally self-limiting. There has been a report of an elevated phenytoin level after 4 weeks of *Paxil* and phenytoin co-administration, and a report of severe hypotension when *Paxil* was added to chronic metoprolol treatment. **DRUG ABUSE AND DEPENDENCE: Controlled Substance Class:** *Paxil* is not a controlled substance. Evaluate patients carefully for history of drug abuse and observe such patients closely for signs of *Paxil* misuse or abuse (e.g., development of tolerance, incrementations of dose, drug-seeking behavior).

behavior). BRS-PX:L12

**SB** SmithKline Beecham Pharmaceuticals Philadelphia, PA 19101



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# PANIC DISORDER

# DEPRESSION

The symptoms may overlap. but the solution is the same



Lifts depression. Lowers associated anxiety symptoms.

of leas or O nause dry mo sweatin female ge decreased nervousnes Paxil in patie oxidase inhibit contraindicated

common adverse events (incidence er and incidence for Paxil at r placebo) in depression disorder studies include ice, abnormal ejaculation pation, asthenia, s, insomnia, tremor, rders, libido decreased e, impotence and ncomitant use of taking monoamine rs (MAOIs) is

Please see brief summary of prescribing information at the end of this advertisement.

CONTROL

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## PHOTO ESSAY

Autism—one of the few neuropsychiatric illnesses to strike in early childhood—is depicted by the universal image of a young person in a state of social isolation. This issue, dedicated to an emerging consensus in autism, includes first-time presentations in the field.



The International Journal of Neuropsychiatric Medicine

Volume 3 • Number 3 March 1998

## **CNS SPECTRUMS**

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Three-pointer

Three Indications. One Product. ZOLOFT.

- Major Depression
  Obsessive-Compulsive Disorder
- 9 Panic Disorder

Please see brief summary of prescribing information on adjacent page. TL155A97

ZOLOFT<sup>®</sup> (sertraline HCl) is indicated for the treatment of depression, obsessions and compulsions in patients with obsessive-compulsive disorder (OCD), and panic disorder. The most common side effects in depression, OCD, and other premarketing controlled trials are nausea, insomnia, diarrhea, ejaculation failure (primarily ejaculatory delay), somnolence, tremor, dyspepsia, increased sweating, anorexia, and decreased libido. The most common side effects in panic disorder trials are diarrhea, ejaculation failure (primarily ejaculatory delay), decreased libido, constipation, anorexia, agitation, tremor, and increased sweating. ZOLOFT is available in 25 mg, 50 mg, and 100 mg scored tablets.

#### BRIEF SUMMARY. Consult the package insert for complete prescribing information

CONTRAINDICATIONS: Concomitant use in patients taking monoamine axidase inhibitors (MAOIs) is contraindicated. WARNINGS: Cases of Construction of a drug interaction between an SRI and an MAOI include: hyperthermia, rigidity, myodanus, autonomic instability toms of a drug interaction between an SSRI and an MAOI include: hyperthermia, rigidity, myodonus, autonomic instability with possible read fluctuations of vital signs, mental status changes that indude contusion, irritability, and extreme ogita-tion progressing to delirium and come. These reactions have also been reported in patients who have recently discontinued an SSRI and have been started on an MAOI. Therefore, it is recommended that ZOLOFT not be used in combination with an MAOI or when the delirium and come. cm SSK1 and have been started on an MAUL. Inereters, it is recommonded mar ZULOPT not be used in commonnon wrm an MAOL, or within 14 days of discontinuing treatment with an MAOL. Similarly, or loss 14 days should be allowed after stopping ZOLOFT before starting an MAOL. PRECAUTIONS: General—Activation of Mania/Hypomania – During premarketing testing, hypomania or mania occurred in approximately 0.4% of ZOLOFT treated patients. Activation of mania/hypomania has also be n reported in a small proportion of patients with Major Affective Disorder treated with other marketed antidepressant and antiobsessional drugs. Weight Loss - Significant weight loss may be an undestrable result of treatment with sentraline for some patients, but an average, patients in controlled trids had minimal. I to 2 pound weight loss, versus smaller changes on placebo. Only rarely have sentraline patients been discontinued for weight loss. **Seizure** – ZOLOFT has not been evaluated in patients with a seizure disorder. These patients were excluded from clinical studies during the product's premarket testing. No seizures were observed among approximately 3000 patients treated with ZOLOFT in the development program for depression. However, 4 patients out of approximately 1800 exposed during the development program for obsessive compulsive disorder experienced seizures, representing a crude incidence of 0.2%. Three of these patients were adolescents, two with a seizure disorder and one with a family history of seizure disorder, none of whom were receiving anticonvulsant medication. Accordingly, ZOLOFT should be introduced with care in patients with a seizure disorder. Suicide - The possibility of a suicide attempt is inherent in depression and may persist until significant remission occurs. Close supervision of high risk patients should accompony initial drug therapy. Prescriptions for ZOLOFT should be written for the smallest auantity of tablets consistent with apod patient management, in order to reduce the risk of overdose. Because of the well-established cornorbidity between both OCD and depression and panic disorder and depression, the same precau tions observed when treating patients with depression should be observed when nearing patients with OCD or point disorder. Weak Uricosuric Effect \_ZOLOFT is associated with a mean decrease in serum unic acid of approximately 7%. The clinical significance of this weak unicosuric effect is unknown, and there have been no reports of acute renal failure with ZOLOFT. Use in Perients with Concomitant Illness — Clinical experience with ZOLOFT in patients with certain concomitant systemic illness is limited. Caution is advisable in using ZOLOFT in patients with diseases or conditions that could affect metabolism or hemodynamic responses. Patients with these diagnoses were excluded from clinical studies during the product's premarket testing. How ever, the electrocardiograms of 774 patients who received ZOLOFT in double-blind trials were evaluated and the data indicate that ZOLOFT is not associate ed with the development of significant ECG abnormalities. ZOLOFT is extensively metabolized by the liver. In subjects with mild, stable cirrhosis of the liver, the clearance of sertraline was decreased, thus increasing the elimination half-life. A lower or less frequent dose should be used in patients with cirrhosis Since 20LOFT is extensively metabolized, excretion of unchanged drug in urine is a minor route of elimination. Until the pharmacokinetics of 20LOFT have been studied in patients with renal impoirment and until adequate numbers of patients with severe renal impoirment have been evaluated during chronic ent with ZOLOFT, it should be used with coution in such potients. Interference with Cognitive and Motor Performance – In controlled studies, ZOLOFF did not cause sedation and did not interfere with psychomotor performance. Hypenatremia – Several cases of hyponatremia have been reported and appeared to be reversible when ZOLOFF was discontinued. Some cases were possibly due to the syndrome of inappropriate antidurence of the severation of the syndrome of the severation of the seve hormone secretion. The majority of these occurrences have been in elderly individuals, some in patients taking diuretics or who were otherwise volume depleted. Platelet Function - There have been rare reports of altered platelet function and/or abnormal results from laboratory studies in patients taking 20.0FL While there have been reports of adnormal bleading or purpura in several patients taking 20.0FL it is unclear whether 20.0FL had a causative role. Information for Patients: Physicians are advised to discuss the following issues with patients for whom they prescribe 20.0FL Patients should be told that although ZOLOFT has not been shown to impair the ability of normal subjects to perform tasks requiring complex motor and mental skills in laboratory experiments, drugs that oct upon the central nervous system may affect some individuals adversely. Patients should be told that although ZOLOFT has not been shown in experiments with normal subjects to increase the mental and motor skill impairments caused by alcohol, the concomitant use of ZOLOFT and alcohol is not advised. Patients should be told that while no adverse interaction of ZOLOFT with over-the-counter (OTC) drug products is known to occur, the potential for interaction exists. Thus, the use of any OTC product should be initiated cautiously according to the directions of products is known to dc.org mit pointent is should be obtained to notify their physician if they be come pregnant or to there to be come pregnant or to there to be come pregnant or to there to be come pregnant or the top there are there are pregnant or top there are there are there are there are there are there are top there are there are top the are top there are top there are top the are top there are top the are top the are top there are top the a drugs. In a study comparing prothrombin time AUC (0-120 hr) following dosing with warfarin (0.75 mg/kg) before and after 21 days of dosing with either ZOLOFT (50-200 mg/day) or placebo, there was a mean increase in prothrombin time of 8% relative to baseline for ZOLOFT compared to a 1% decrease for placebo (p<0.02). The normalization of prothrombin time for the ZOLOFT group was delayed compared to the placebo group. The clinical sig nificance of this change is unknown. Accordingly, prothrombin time should be carefully monitored when ZOLOFT therapy is initiated or stopped. dine - In a study assessing disposition of ZOLOFT (100 mg) on the second of 8 days of cimetidine administration (800 mg daily), there were significant There in a study cases and application of court if the migration of a study is a diministrational (court may), mere very significant increases in (200F) mean AUC (CoHT) (court if and harifiel (2005) compared to the placebag group. The clinical significance of these changes is unknown. **CNS Active Drugs** — In a study comparing the disposition of introvenously administered diazepum before and offer 21 days of dosing with either 200FT (SD to 200 mg/day escalating dose) or placebo, there was a 23% decrease relative to baseline in diazepum cheanance for the 200FT group compared to a 19% decrease relative to baseline for the placebo group (p-c0.03). There was a 23% increase in Tranx for desmethyldizaepum in the 200FT group compared to a 19% decrease in the juncebo group (p-c0.03). The dinical significance of these changes is unknown. In a placebo-controlled trial in normal volumeers, the doministration of two doses of 200FT did not significantly after steady-state lithium levels or the renal clearance of lithium. Nonetheless, at this time, it is recommended that plasma lithium levels be monitored following initiation of ZOLOFT therapy with appropriate adjustments The hillitum dase. The risk of using 2010FT in combination with other CNS active drugs has not been systematically evaluated. Consequently, curdion is advised if the concomitant administration of 2010FT and such drugs is required. There is limited controlled experience regarding the optimal timing of switching from other antidepressons to 2010FT. Care and prudent medical judgment should be exercised when switching, particularly from long-acting agents. The duration of an appropriate washour period which should intervene before switching from one selective serotowin reuptake inhibitor (SSRI) to another has not been established. Drugs Metabolized by P450 3A4 — In two separate in vivo interaction studies, settraline was coadministered with the cytechrome P450 3A4 substrates, terfenadine or corbamozepine, under steady-state conditions. The results of these studies demonstrated that sertraline coordinistration did not increase plasma concentrations of terfenadine or carbamazepine. These data suggest that sertraline's extent of inhibition of P450 3A4 activity is not likely to be of clinical significance. Drugs Metabolized by P450 2D6 - Many antidepressants, e.g., the SSRIs, including sentraline, and most tricyclic antidepressants inhibit the biachemical activity of the drug metabolizing isozyme cytochrome P450 206 (debrisa-quin hydroxylase), and, thus, may increase the plasma concentrations of coadministered drugs that are metabolized by P450 206. The drugs for which this potential interaction is of greatest concern are those metabolized primarily by 2D6 and which have a narrow therapeutic index, e.g., the tricyclic anidepressants and the Type 1C antianthythmics proparement of flecalinide. The extent to which this interaction is an important clinical problem depends on the extent of the inhibition of P450 2D6 by the antidepressant and the therapeutic index of the coordiministered drug. There is variability among the antidepressants in the extent of clinically important 2D6 inhibition, and in fact sertrafine at lower doese has a less prominent inhibitory effect on 2D6 than amenometrics in the doss. Neverthese, even sentration has the potential for clinically important 2005 inhibition. Consequently, concomitant use of a drug metabolized by P450 206 with ZOLOFT may require lower doss than usally prescribed for the other drug. Furthermore, whenever ZOLOFT is withdrawn from co-therapy, an increased doss of the coadministered drug may be required (see Tricyclic Antidepressants under PRECAUTIONS). **Fricyclic Antide**pressants (TCAs) — The extent to which SSRI-ICA interactions may pase clinical problems will depend on the degree of inhibition and the pharma-cokinetics of the SSRI involved. Nevertheless, caution is indicated in the coadministration of TCAs with ZOLOFT, because settraline may inhibit TCA metabolism. Plasma TCA concentrations may need to be monitored, and the dose of TCA may need to be reduced, if a TCA is co-administered with ZOLOFT Insulation that it is a second sec tolbutamide following an intravenous 1000 mg dose. ZOLOFT administration did not noticeably change either the plasma protein binding or the apparent volume of distribution of tollautamide, suggesting that the decreased clearance was due to a change in the metabolism of the drug. The clinical significance of this decrease in talbutamide clearance is unknown. Atenalal - ZOLOFT (100 mg) when administered to 10 healthy male subjects had no effect an the betroadrenergic blocking ability of atenolol. Digoxin — In a placebo-controlled trial in normal volunteers, administration of ZOLDFT for 17 days (including 200 mg/day for the last 10 days) did not change serum digoxin levels or digoxin renal clearance. Microsomal Enzyme Induction — Preclinical studies have shown ZOLOFT to induce hepatic microsomal enzymes. In clinical studies ZOLOFT was shown to induce hepatic enzymes minimally as determined by a small (5%) but statistically significant decrease in antipyrine half-life following administration of 200 mg/day for 21 days. This small change in antipyrine holf-life reflects a clinically insignificant change in hepatic metabolism. **Electroconvolsive Therapy** — There are no clinical stud-ies establishing the risks or benefits of the combined use of electroconvolsive therapy (ECT) and ZOLOFT. **Alcohol** — Although ZOLOFT did not potentiate the cognitive and psychomotor effects of alcohol in experiments with normal subjects, the concomitant use of ZOLOFT and alcohol is not recommended. Carcinogenesis, Mutagenesis, Impairment of Fertility: Lifetime carcinogenicity studies were carried out in CD-1 mice and Long-Evans rats at

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doese up to 40 mg/kg/day. These doese correspond to 1 finnes (mice) and 2 finnes (rats) the maximum recommended human dose (MRHb) on a mg/m<sup>2</sup> basis. There was a doserelated increase of liver adenomas in male mice receiving sentaline at 10-40 mg/kg (0.25 - 1.0 finnes the MRHD on a mg/m<sup>2</sup> basis). No increase was seen in fendle mice or in rats of either sex receiving sentaline at 10-40 mg/kg (0.25 - 1.0 finnes the MRHD on a mg/m<sup>2</sup> basis). No increase was seen in fendle mice or in rats of either sex receiving sentaline at 10-40 mg/kg (0.25 - 1.0 finnes the MRHD on a mg/m<sup>2</sup> basis). No increase was seen in fendle mice or in rats of either sex receiving sentaline at 10-40 mg/kg (0.25 - 1.0 finnes the MRHD on a mg/m<sup>2</sup> basis). The second of the thyroid in female rats receiving sentaline at 40 mg/kg (2 finnes the MRHD on a mg/m<sup>2</sup> basis); organeed to placeba controls, this effect more and early durknown significance to humans. There was an increase in follicular adenomas there a varies an increase in teerine adenocarcinomas in rats receiving sentaline at 10-40 mg/kg (0.5 - 2.0 finnes the MRHD on a mg/m<sup>2</sup> basis); compared to placeba controls, this effect was not dearly drug related. Sentaline had no genotaxis effect, with ar without methodik activation, based on the following assays: bacterial mutation assay, mouse hymphorne mutation assay, and tests for cytogenetic aberrations in wive in mouse bane marrow and in withe in human hymphocytes. A decrease in femility was seen in one of two rat studes at a does of 80 mg/kg (4 finnes the maximum human dose on a mg/m<sup>2</sup> basis). **Pregnancy — Pregnancy Caregory C** There are no adequate and well-controlled studies in pregnant women. **200**(F1 should be used during pregnancy only if the potential barlifies the potential tisk to the fetus. **Labor and Delivery** — In effect of 2010FT on labor and delivery in humans is unknown. **Muesing Mothers —** It is not known whether, and fs in what amount, sentaline or its metabolites are excreted in human milk, caution should be

- 1	NOST COMMON	TREATMENT	-EMERGENT AC	VERSE EVENT:	5: INCIDENCE IN PLACEBO	-CONTROLLED CLINICAL TRIALS

	PERCENTAGE OF PATIENTS REPORTING EVENT					
	Depression/Other*		OCD		Panic D	lisorder
BODY SYSTEM/ ADVERSE EVENT	ZOLOFT (N=861)	Placebo (N=853)	ZOLOFT (N=533)	Placebo (N=373)	ZOLOFT (N=430)	<b>Placebo</b> (N=275)
Autonomic Nervous System Disorders						
Ejaculation Failure (1)	7	<1 3	17	2	19	1
Sweating Increased	8	3	6	1	5	1
Central & Peripheral Nervous System Disorders						
Somnolence	13	6	15 8	8	15 5	9
Tremor	11	3	8	1	5	
Gastrointestinal Disorders						
Anorexia	3	2	11	2	7	23
Constipation	8	6	6	4	7	3
Diarrhea/Loose Stools	18	9	24 10	10	20	9
Dyspepsia	6	3 12	10	4	10	8 18
Nausea	26	12	30	11	29	18
Psychiatric Disorders						
Agitation	6	4	6	3	6	2
Insomnia	16	9	28	12	25	2 18
Libido Decreased	1	<1	11	2	7	1
(1) Bin site simulation delay. Description and use for male actions only (1) 271 2010(T description (attacks 1) 271 placebo						

(1)Primarily ejeculatory delay. Denominator used was for male patients only (N=271 ZOLOFT depression/other\*; N=271 placebo depression/other\*; N=296 ZOLOFT OCD; N=219 placebo OCD; N=216 ZOLOFT panic disorder; N=134 placebo panic disorder). \*Depression and other premarketing controlled trials.

Associated With Discontinuation of Treatment: The adverse events associated with discontinuation of ZOLOFT treatment (incidence at least twice that for placebo and at least 1% for ZOLOFT ) in depression and other premarketing controlled trials are agitation, diarrhea, dry mouth, ejaculation failure (primarily ejaculatory delay), headache, insomnia, nausea, somnolence, and tremor; in OCD are diarrhea, dizziness, ejaculation failure (primarily ejaculatory delay), insomnia, nausea, and somnolence; in panic disorder are agitation, anorexia, anxiety, impaired concentration, depersonalization, diarthea, dizzines, dry moth, dyspassie, ejeculation failure (primarily ejaculatory delay), fatigue, headache, insomnia, nausen, nervousness, paresthesia, somnolence, and vaniting. Other Events Observed Daving the Premarketing Evaluation of ZOLOFF: During its premarketing assess ment, multiple doses of ZOLOFF were administered to opproximately 3800 subjects. Events are further categorized by body system and listed in order of decreasing frequency according to the following definitions: frequent adverse events are those accurring on one or more accasions in at least 1/100 patients (only those not already listed in the tabulated results from placebo-controlled trials appear in this listing); infrequent adverse events are those occurring in 1/100 to 1/1000 patients; rare events are those occurring in fewer than 1/1000 patients. Events of major clinical importance are also ed in the PRECAUTIONS section. Autonomic Nervous System Disorders - Frequent: impotence; Infrequent: flushing, increased saliva, cold damme in Kachina section section in the property of the section of the se hypertension, myocardial infarction, cerebrovascular disorder. Central and Peripheral Nervous System Disorders - Frequent: hyper hypoesthesia; Infrequent: twitching, confusion, hyperkinesia, vertigo, ataxia, migraine, abnormal coordination, hyperesthesia, leg cramps, abnormal gait, nystogmus, hypokinesia; Rare: dysphonia, coma, dyskinesia, hypotonia, ptosis, choreoarthetasis, hyporeflexia. Disorders of Skin and Appendages — infrequent: pruritus, acne, unicario, alopecia, dry skin, erythematous rash, photosensitivity teaction, maculopapular rash; Rare: folicalar rash, eczerna, dermatitis, contact dermatitis, bullous eruption, hypertrichosis, skin discoloration, pustular rash. Endocrine Disorders - Rare: exophan nam, so secondo contra contr tum hemorrhage, hemorrhagic peptic uker, proctitis, ulcerative stomatitis, tongue edema, tongue ulceration. General - Frequent: back pain, asthenia, malaise, weight increase, Infrequent: fever, rigors, generalized edema; Rare: face edema, aphthous stamatitis. Hearing and Vestibuler Disorders — Rare: hyperocusis, labyrinthine disorder. Hematopoletic and Lymphatic — Rare: anemia, anterior chamber eye hemorrhage. Liver and Billiary System Disorders — Rare: abnormal hepatic function. Metabolic and Nutritional Disorders — Infrequent: thirst; Rare: hypoglycemia, hypo glycemia reaction. Musculoskeletal System Disorders — Frequent: mydaja; Infrequent: arthvalaja, dystania, arthvosis, muscle aramps, muscle weakness. Psychiatric Disorders — Frequent: yawning, other male sexual dysfunction, other female sexual dysfunction; Infrequent depression, amnesia, paronina, teeth-grinding, emotional lability, apathy, abnormal dreams, euphonia, parancid reaction, halkucination, aggressive reaction, aggrevatsion, delusions; Rare: withdrawal syndrome, suicide ideation, libido increased, somnambulism, illusion. Reproductive - Infrequent: men stradi disorder, dysmenorthea, intermenstradi bleeding, voginal hemorthage, anenorthea, leukorthea; Rare: ferrale breast pain, menorthagia, bal-anaposthitis, breast enlargement, atrophic voginitis, acute ferrale mastitis. Respiratory System Disorders — Frequent, thinitis; Infrequent; coughing, dyspnea, upper respiratory tract infection, epistaxis, branchospasm, sinusitis; Rare: hyperventilation, bradypnea, stitidor, apnea, branchitis, hemoptysis, Upporentificion, laryngismus, laryngistus, Special Senses – Frequent tinnitus, infraentament conjunctivitis, grandie, eye pain, abnormal accommodation, Rare: xerophthalmia, photophobia, diplopia, abnormal locrimation, scotoma, visual field defect. Urinary System Disorders – Infraquent: micturition frequency, polyuria, urinary retention, dysuria, nocturia, urinary incontinence; Rare: cystitis, oliguria, pyelonephritis, hematuria, renal pain, strangury. Laboratory Tests: In man, asymptomatic elevations in serum transaminases (SGOT [or AST] and SGPT [or ALT]) have been reported infrequently (approximately 0.8%) in association with ZOLOFT administration. These hepatic enzyme elevations usually occured within the first 1 to 9 weeks of drug treatment industry of an insistantian with 2007 rounningstantary. These points exprise elevations baday occured within the first in 27 weeks to day inclusion and promptly diministrated upon day discontinuation. 200,017 therapy was associated with small mean increases in total cholesterol (approximately 3%) and highyerides (approximately 5%), and a small mean decrease in serum unic add (approximately 7%) of no apparent dinical importance. The safety profile abserved with 200,017 treatment in patients with depression, OCD and point disorder is similar. **Other Events Observed During the Postmarket** ing Evaluation of ZOLOFT - Reports of adverse events temporally associated with ZOLOFT that have been received since market introduction, that are and extraction of external intervences of the second s severe cutaneous disorders, rare reports of pancreatitis, and liver events — clinical features (which in the majority of cases appeared to be reversible with dis-continuation of 20L0FI) occurring in one or more patients include: elevated enzymes, increased bilitubin, hepatomegady, hepatitis, jaundice, abdominal pain, vomiting, liver failure and death. **OVERDOSAGE:** Symptoms of overdose with 20L0FT alone included somnolence, nausea, vomiting, pany, terming, the name and additional pupils. Featment was primarily supporte and included manifolding and use of activated duracid, gastric large or catherits and hydration. Although there were no reports of death when ZOLOFT was taken alone, there were 4 deaths involving overdoses of ZOLOFT in combination with other drugs and/or alcohol, as of November 1992. Therefore, any overdosage should be treated aggressively.



THE SEAVER AUTISM RESEARCH CENTER at The Mount Sinai School of Medicine, in association with the AUTISM SOCIETY OF AMERICA and CURE AUTISM NOW, present a full day symposium:

# New Insights in the Diagnosis, Neurobiology, Genetics and Treatment of Autism Saturday, May 2, 1998, Mount Sinai Medical Center, New York City

A partial list of speakers include: Eric Hollander, MD, Joseph Buxbaum, MD, Gina DelGiudice-Asch, MD, Charles Cartwright, MD (Mount Sinai School of Medicine); Mr. Hirschell E. Levine (Beatrice and Samuel A. Seaver Foundation); John Maltby (Autism Society of America); Portia Iversen (Cure Autism Now); Clarence Schutt, PhD (National Alliance for Autism Research); Elizabeth Leonard, PhD (Barrow Neurological Institute); Robert Schultz, PhD (Yale University); Guest Speaker-Temple Grandin, PhD (Colorado State University)

A partial list of topics include: Research Update: Seaver Autism Research Center; Molecular Biology and Genetics of Autism; Pharmacological Treatment of Autism; Functional Imaging Studies of Autism; Autoimmune Function in Autism; Psychopharmacology Workshop; Consumer Advocacy Workshop; Social Skills Treatment Workshop



## **REGISTRATION FORM**

Symposium on New Insights in Diagnosis, Neurobiology, Genetics and Treatment of Autism, Saturday, May 2, 1998, Mount Sinai Medical Center (Hatch Auditorium), New York

Name		N	
Address			
City	State	Zip	
Daytime Telephone			
Tuition Fees: Physicians and Other Health Prof Mount Sinai Alumni Associates . Fellows and Residents Family Members and Consumers			\$100.00 .\$75.00
Please mail your check with regi The Page and William Black Post Box 1193			
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