LUVOX (fluvoxamine maleate) 25 mg TABLETS, 50 mg and 100 mg SCORED TABLETS

Brief Summary (For full Prescribing Information and Patient Information, refer to package insert.)

INDICATIONS AND USAGE

LUVOX* Tablets are indicated for the treatment Disorder (OCD), as defined in the DSM-III-R. tment of obsessions and compulsions in adults and children and adolescents (ages 8-17) with Obsessive Compulsive

CONTRAINDICATIONS

Co-administration of terfenadine, astemizole, or cisapride with LUVOX® Tablets is contraindicated (see WARNINGS and PRECAUTIONS).

LUVOX® Tablets are contraindicated in patients with a history of hypersensitivity to fluvoxamine maler

TRANSINGS
In patients receiving another serotonin reuptake inhibitor drug in combination with monoamine oxidase inhibitors (MAOI), there have been reports of serious, sometimes fatal, reactions. Some cases presented with features resembling neuroleptic madignant syndrome. Therefore, it is recommended that LUVOX" Tablets not be used in combination with a MAOI, or within 14 days of discontinuing treatment with a MAOI. After stopping LUVOX" Tablets, at least 2 weeks should be allowed before

14 adys or discontinuing treatment with a MAOI. After stopping LUVUA" tablets, at least x weeks should be allowed before starting a MAOI.

Terfenadine, astemizole and cisapride are all metabolized by the cytochrome P450IIIA4 isoenzyme. Increased plasma concentrations of terfenadine, astemizole and cisapride cause QT prolongation and have been associated with torsades de pointes-type ventricular tachycardia, sometimes fatal. Although it has not been definitively demonstrated that fluvoxamine is a potent IIIA4 inhibitor, it is likely to be. Consequently, it is recommended that fluvoxamine not be used in combination with either terfenadine, astemizole, or cisapride.

with either terfenadine, astemizole, or cisapride.

Other Potentially Important Drug Interactions
(Also see PRECURIOMIS) - Drug Interactions
(Also see PRECURIOMIS) - Drug Interactions (Programmer of these drugs is likely to be reduced by hepotic oxidation (e.g., alprazolam, midazolam, triazolam, etc.) should be used with courton because the clearance of these drugs is likely to be reduced by Havozamine. The clearance of benerodizergom, ternazopam) is unlikely to be riflected by Havozamine. Apprazolam: When filavozamine maleate (100 mg qd) and alprazolam (1 mg qid) were co-administered to steady state, plasma concentrations and other pharmocokinetic parameters (AUC, C..., T.) of alprazolam were opproximately twice those observed when alprazolam was administered alone; and clearance was reduced by about 50%. The elevated plasma alprazolam concentrations resulted in decreased psychomotra performance and memory. This interaction, which has not been investigated using higher doese of throwasmine, may be more pronounced if a 300 mg daily does is co-administered, particularly since fluvoxomine exhibits non-linear pharmacokinetics over the disage range 100-300 mg. It algoractom is co-administered with LUVOX" flobles, the initial alprazolam disages that the conditional function to the lowest effective does is recommended. No dosage adjustment is requised for LUVOX" flobles. Diazagema: The co-administration of LUVOX" flobles and diazagema is a strong its elevation of LUVOX" flobles. Diazagema: The co-administration of LUVOX" flobles and diazagema is a strong its elevation of the strong through the co-administration of LUVOX" flobles. Diazagema: The effect of steady-state fluvoxamine (50 mg bid) on the pharmacokinetics of a single dose of theophylline (375 mg as 442 mg antimophylline) was evoluted in 12 healthy non-smoking, male volunteers. The clearance of theophylline was decreased approximately 34od. Therefore, if theophylline is ordinistration melacute, its does should be reduced to one third of the waste does and plasma contentrations of theophylline should be monitored. No dosage adjustment is required for LUVOX" flobles. Warfarine. When fluvoxamine melacute (50 mg tid) was patients receiving and anticogulants and LUVOX" flobles should have their protrioration increased by 98% and protrioration intrinsesser protriorally. No dosage adjustment is required for LUVOX" flobles. Core. T.; of alprazolam were approximately twice those observed when alprazolam was administered alone; oral clearance was reduced by about 50%. The

PRECAUTIONS

General

General

Activation of Mania/Hypomania: During premarketing studies involving primarily depressed patients, hypomania or mania occurred in approximately 1% of patients treated with filtrovamine. Activation of mania/hypomania has also been reported in a small proportion of patients with mojor affective disorder who were treated with other marketed antidepressents. As with all antidegressants, LIVOX** Tablets should be used cautiously in patients with a history of mania. Seizures: During premarketing studies, seizures were reported in 0.2% of flovoxomine-treated patients. LIVOX** Tablets should be associated in any patient who develops seizures. Seizures were largered in any patient who develops seizures. Seizures were largered in any patient who develops seizures. Seizures were maintained in any patient who develops seizures. Seizures were maintained in any patient who develops seizures. Seizures were maintained in any patient who develops seizures. Seizures vollets in solicited and the patients is inherent in patients with allowers with a patients with consistent with good patient management in order to reduce the risk of overdose. Use in Patients with Concentiant Illness: Closely monitored clinical experience with LIVOX** Tablets in patients with concentiant systemic illness is limited. Coution is advised in administrating LIVOX** Tablets to patients with developed in administrating LIVOX** Tablets in patients with concentiant patients with concentiant systemic illness is limited. Coution is advised in administrating LIVOX** Tablets in patients with the developed from many clinical studies advised in promotering tenders between fluvoxamine and placebo in the emergence of clinically important ECG changes. In patients with their dysfunction, fluvoxamine clearance was decreased by approximately 30%. LIVOX** Tablets should be slowly introd in patients with her dysfunction of intention of treatment. titrated in patients with liver dysfunction during the initiation of treatment.

Information for Patients

Information for Patients

Physicians are advised to discuss the following issues with patients for whom they prescribe LUVOX** Toblets: Interference with Cognitive or Motor

Performance: Since any psychocitive drug may impair judgement, thinking, or motor skills, patients should be coutioned about operating hazardous
machinery, including automobiles, until they are certain that LUVOX** Toblets therapy does not adversely affect their ability to engage in such activities.

Pergenancy: Patients should be obvised to notify their physicians if they become pregnant or intend to become pregnant during therapy with LUVOX**
Toblets. Nursing: Patients receiving LUVOX** Toblets should be advised to notify their physicians if they are braces feeding an infant. (See PRECAUTIONS

- Nursing Mothers.). Concomitant Medications: Patients should be advised to notify their physicians if they are bracing, or plan to take, any prescribing in protent interactions with LUVOX**, Toblets. Allored: As with other systychropic medications, patients should be advised to notify their physicians if they develop a rosh, hives, or a related allergic phenomenon during therapy with LUVOX** Toblets.

There are no specific laboratory tests recommended.

Laboratory Tests
There are no specific blooratory tests recommended.

Drug Interactions

Potential interactions with drugs that inhibit or are Metabolized by Cytochrome P450 Isozymes: Multiple hepatic cytochrome P450 (CY450) enzymes are involved in the oxidative biotransformation of a large number of structurally different drugs and endogenous compounds. The oxidable knowledge concerning the relationship of fluoroxamine and the CYP450 enzyme system has been obtained mostly from phormocokinetic interlock studies conducted in healthy voluntees, but some preliminary in vitro data are also available. Based on a finding of substantial interactions of fluoroxamine with certain of these and limited in vitro data for the IIIA4 isoenzyme, it appears that fluoroxamine inhibits isoenzymes that are known to be involved in the metabolism of drugs such as wardning, theophyline and programolal. A clinically significant fluoroxamine interaction is possible with drugs having a narrow therapeutic mixed such as teteraction, and drug that is eliminated via oxidative metabolism and has a narrow therapeutic window, plasma levels and/or pharmocodymate effects of the later drug should be monitored closely, or less studies and an arrow therapeutic window, plasma levels and/or pharmocodymate effects of the later drug should be monitored closely, or less studies and an arrow therapeutic window, plasma levels and/or pharmocodymate effects of the later drugs should be monitored closely, or less studies and the studies of the studies and the studies and

daily dose on a mg/m² basis) had no effect on mating performance, duration of gestation, or pregnancy rate.

Pregnancy
Tearlogenic Effects: Pregnancy Category C: In tentology studies in rats and robbits, daily and doses of fluvoxamine maleate of up to 80 and
40 mg/kg, respectively (approximately 2 himes the maximum human daily dose on a mg/m² basis) caused no fetal malformations. However, in other
reproduction studies in which pregnant rats were dosed through wearing there was (1) on increase in pup mortality at birth (seen at 80 mg/kg), and (2) decreases in postnatula pup weights (seen at 160 but not at 80 mg/kg) and survival (seen at all doses; lowest dose tested
= 5 mg/kg). Obese of 5, 20, 80, and 160 mg/kg are approximately 0.1, 0.5, 2, and 4 times the maximum human daily dose on a mg/m² basis.)
While the results of a cross-fostering study implied that at least some of these results likely occurred secondarily to maternal toxicity, the role of a direct drug effect on the fetuses or purs contains and a major and the second of the second o

Nursing Mothers

NOTION IN MONEY.

As for many other drugs, fluvoxomine is secreted in human breast milk. The decision of whether to discontinue nursing or to discontinue the drug should take into account the potential for serious adverse effects from exposure to fluvoxomine in the nursing infant as well as the potential benefits of LUYOX** (fluvoxomine maleate) Toblets therapy to the mother.

Predicting of fluoroamine molecte for the treatment of Obsessive Compulsive Disorder was demonstrated in a 10-week multicenter placeba controlled study with 120 outpatients ages 8-17. The adverse event profile observed in that study was generally similar to that observed in adult studies with fluoroxamine (see ADVERSE REACTIONS and DOSAGE AND ADMINISTRATION).

Decreased appetite and weight loss have been observed in association with the use of fluvoxomine as well as other SSRIs. Consequently, regular monitoring of weight and growth is recommended if treatment of a child with an SSRI is to be continued long term.

Geriatric Use

Approximately 200 parients participating in controlled premarketing studies with LUVOX[®] Tablets were 65 years of age or over. No overall differences in safety were abserved between these patients and younger patients. Other reported clinical experience has not identified differences in response between the elderly and younger patients. However, the clearance of fluvoxamine is decreased by about 50% in elderly compared to younger patients (see Pharmacokinetics under CLINICAL

PHARMACOLOGY), and greater sensitivity of some older individuals also cannot be ruled out. Consequently, LUVOX® Tablets should be slowly titrated during initiation

Associated with Discontinuation of Treatment

Of the 1087 OCD and depressed patients treated with fluvoxamine maleate in controlled clinical trials conducted in North America, 22% discontinued treatment due to an adverse event.

Incidence in Controlled Trials - Commonly Observed Adverse Events in Controlled Clinical Trials: LUVOX® Tablets have been studied

Incidence in Controlled Trials - Commonly Observed Adverse Events in Controlled Clinical Trials: LIVOX® Toblets have been studied in controlled that of COL (18-320) and depression (18-1350). In general, otherse event rates were similar in the two data sets as well as in the pediate (105 study, the most commonly observed adverse events associated with the use of LIVOX® Toblets and likely to be duryelated (incidence of 5% or genete and at least twice that for placebo) derived from Toble 1 were: somnolence, insomnia, nervousness, trema, nousea, dyspepsia, anarexia, vamiting, abnormal ejaculation, asthemia, and sweating. In a pool of two stables involving only patients with OCD, the following additional events were identified using the above rule: dry nount, decreased libida, unionary frequency, anagoram, chimitis and larts perversion. In a study of pediatric patients with OCD, the following additional events were identified using the above rule: agitation, depression, dysmenorthea, flatulence, hyperkinesia, and rash.

Adverse Events Occurring at an Incidence of 1%: Toble 1 enumerates adverse events that occurred at to frequency of 1% or more, and were more frequent than in the placebo group, among potients treated with LIVOX® Tobles in two short-term placebo controlled OCD trials (10 week) and depression trials (6 week) in which patients were dosed in a range of generally 100 to 300 mg/day. This table shows the percentage of patients in each group who had at least one occurrence of an event at some time during their teatment. Reported oberse events were classified using a standard COSTART-based Dictionary terminology, the prescriber should be aware that these figures cannot be used to predict the incidence of side effects in the cause of usual medical practice where patient characteristics and other fortors may differ from those that prevailed in the clinical trials. Similarly, the cited frequencies cannot be used to predict the incidence of side effects in the cause of usual medical practice where potient charact do provide the prescribing physician with some basis for estimating the relative contribution of drug and non-drug factors to the side-effect incidence rate in

me populations studies. Tracament-emergent adverse event incidence rates by Body System in adult ocd and depression Populations Combined ((hyxocomine (N=892) vs. plocebo (N=778) by potents-percentage): Body As Whole: Heodoche (22 vs. 20); Asthenia (14 vs. 6); Flu Syndrome (3 vs. 2); Chills (2 vs. 1). Cardiovascular: Polpitations (3 vs. 2). Digestive system: Nousea (40 vs. 14); Durcheo (11 vs. 7); Constipation (10 vs. 8); Dyspepia (10 vs. 5); Anaexia (6 vs. 2); Voloniting (5 vs. 2); Flotulence (4 vs. 3); Toolh Disorder (3 vs. 1); Dysphagia (2 vs. 1), NERVOUS SYSTEM: Sormolence (22 vs. 8); Insonnia (21 vs. 10); Dry Mouth (14 vs. 10); Nervousness (12 vs. 5); Dizziness (11

Dysphogia (2 vs. 1). NERVOUS SYSTEM: Someleires (22 vs. 8); Insomini (21 vs. 10); Bry Mouth (14 vs. 10); Nerousness (12 vs. 5); Diszines (17 vs. 6); Itemor (5 vs. 1); Anxiety (5 vs. 3); Vissodilatotion" (3 vs. 1) Hypertonia (2 vs. 1), Agliation (2 vs. 1); Decreased Libido (2 vs. 1); Depression (2 vs. 1); Simulation (2 vs. 1); RSSImulation (2 vs. 1); Anxiety (3 vs. 1); Anxiety (3 vs. 2); UROGENITAL: Abnormal Epoulation. A (8 vs. 1); Libinary Frequency (3 vs. 2); Important (2 vs. 1); Anxiety (3 vs. 2); Important (3 vs. 2); Important (4 vs. 1); Anxiety (4 pharyngitis, agitation, impotence, myoclonus/twitch, thirst, weight loss, leg aramps, myalgia and urinary retention. These events are listed in order of decreasing

Other Adverse Events in OCD Pediatric Population. In Pediatric patients (N=57) heated with LUVOX" Tablets, the overall profile of adverse events is similar to that seen in adult studies. Other nearlines which have been reported in two or more pediatric patients, and were more frequent than in the placebo group group were: abnormal thinking, cough increase, dysmenorthea, ecclymosis, emotional lability, epistosis, hyperkinesia, infection, manic reaction, rash, sinusitis, and

Vital Sign Changes

Comparisons of fluvoxamine maleate and placebo groups in separate pools of short-term OCD and depression trials on (1) median change from baseline on various vital signs variables and on (2) incidence of patients meeting criteria for potentially important changes from baseline on various vital signs variables revealed no important differences between fluvoxamine maleate and placebo.

Leboratory Changes
Comparisons of fluvoxamine malerate and placebo groups in separate pools of short-term OCD and depression triols on (1) median change from baseline on various serum chemistry, hematology, and urinalysis variobles and on (2) incidence of patients meeting criteria for potentially important changes from baseline on various serum chemistry, hematology, and urinalysis variobles revealed no important differences between fluvoxamine malerate and placebo.

besiline on various serum chemistry, hemotology, and uninalysts variousles revealed no important differences between fluvoxamine modeled and placebo.

EGG Changes

Comparisons of fluvoxamine modelete and placebo groups in separate pools of short-term OCD and depression trials on (1) mean change from baseline on various ECG variables and on (2) incidence of patients meeting criteria for potentially important changes from baseline on various ECG variables and on (2) incidence of patients meeting criteria for potentially important changes from baseline on various ECG variables revealed no important differences between fluvoxamine meliated and placebo.

Other Events Observed During the Premarketting Evaluation of LUVOX® Tablets

During premarketing clinical trials conducted in North America and Europe, multiple doses of fluvoxamine maleate were administered for a combined total of 2737 patient exposures in patients suffering OCD or Major Depressive Bisander. Univoxed events associated with this exposure were recorded by clinical investigators using descriptive terminology of their own choosing. Consequently, it is not possible to provide a meaningful estimate of the proportion of individuals experiencing obverse events without first grouping similar types of untrowed events into a limited (i.e., reduced) number of standard event events as a general as to be uninformative; it was replaced with a more informative term. The frequencies presented, therefore, represent the proportion of the 2737 potient exposures to multiple doses of fluvoxamine meladete. We experienced an event of the type cited on all electronic or one occasion within exceeding exceptions. If the common adverse experiences in placebo-controlled OCD and depression clinical trials, are excluded; 21 those events for which a drug cause was considered remote (i.e., neplosia, postivintestical carcinoma, herpes simplex, herpes zosta, application is included. It is important to emphasize that, although the events separated in one or docreasion within thoughts included. It is important to emphasize that, although the events reported did occur during treatment with floworamine moderus, a cuscal relationship to floworamine moderus has not been established. Events or enturher dossfind within body system cropposes and enumerated in order of devent excluding frequency using the following definitions: frequent odverse events are those occurring on one or more occasions in at least 1/100 patients, giving the following definitions: frequent odverse events are those socrating in less than 1/1000 patients, giving the following definitions: frequent odverse events are those socrating in less than 1/1000 patients, giving the following definitions: frequent odverse events are those socrating in less than 1/1000 patients. Body of the control of the following definitions: frequent ordinates of the following definitions or the following definiti

Based on the number of females. Based on the number of males

Non-US Postmarketing Reports

Voluntary reports of adverse events in patients taking LUVOX[®] Tablets that have been received since market introduction and are of unknown causal relationship to LUVOX[®] Tablets use include: toxic epidermal necrolysis, Stevens-Johnson syndrome, Henoch-Schoenlein purpura, bullous eruption, priopism, agranulos/prosis, reunropathy, apolisis, comercia, prophyritic reaction, hyponatremia, acute renal failure, hepatitis, and severe akinesia with fever when fluvoxamine was co-administered with antipsychotic medication.

OVERDOSAGE

Refer to package insert (11E Rev 3/98) for overdosage information. **DOSAGE AND ADMINISTRATION**

Reference: 1. Data on file, Solvay Pharmaceuticals, Inc.

© 1999 Solvay Pharmaceuticals, Inc. All rights reserved.

Refer to package insert (11E Rev 3/98) for dosage and administration information.

Rev 10/98 (11E-5)

Solvay Pharmaceuticals

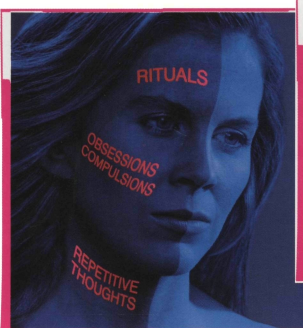
Marietta, GA 30062

Pharmacia&Upjohn

Solvay **Pharmaceuticals** January 1999

OCD IS AN ANXIETY DISORDER

from the profound anxiety of OCD





VISIT THE OCD WEB SITE AT http://www.ocdresource.com

SIGNIFICANTLY IMPROVES OBSESSIVE-COMPULSIVE SYMPTOMS¹

LOW INCIDENCE OF AGITATION IN ADULTS1

▼ 2% vs 1% for placebo

LOW INCIDENCE OF SEXUAL DYSFUNCTION¹

▼ LUVOX® Tablets vs placebo*: decreased libido 2% vs 1%; delayed ejaculation 8% vs 1%; anorgasmia 2% vs 0%; impotence 2% vs 1%

FAVORABLE TOLERABILITY PROFILE¹

- ▼ For adults, the most commonly observed adverse events compared to placebo were somnolence 22% vs 8%; insomnia 21% vs 10%; nervousness 12% vs 5%; nausea 40% vs 14%; asthenia 14% vs 6%
- ▼ Adverse events in children and adolescents were similar to those observed in adult studies. The most commonly observed adverse events compared to placebo were: agitation 12% vs 3%; hyperkinesia 12% vs 3%; depression 5% vs 0%; dysmenorrhea 7% vs 3%; flatulence 5% vs 0%; rash 7% vs 3%
- ▼ Concomitant use of LUVOX® Tablets and monoamine oxidase inhibitors is not recommended
- ▼ Fluvoxamine should not be used in combination with terfenadine, astemizole, or cisapride

^{*}Parameters occurring ≥1% with fluvoxamine maleate.



Please see brief summary of prescribing information on adjacent page.

©1999 Solvay Pharmaceuticals, Inc. All rights reserved.

fluvoxamine maleate 25 mg TABLETS 50 mg & 100 mg SCORED TABLETS

THE #1 SSRI PRESCRIBED BY PSYCHIATRISTS FOR OCD