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

Brief Summary of prescribing information (based on 8E1252 Rev 3/97)

# INDICATIONS AND USAGE

LIVOX Tablest are indicated for the treatment of obsessions and compulsions in patients with Obsessive Compulsive Disorder (OCD), as defined in the DSM-IIR. Obsessive Compulsive Disorder is characterized by recurrent and persistent ideas, thoughts, impulses or images (obsessions) that are ego-dystonic and/or repetitive, purposeful, and intentional behaviors (compulsions) that are recognized by the person as excessive or unreasonable.

# CONTRAINDICATIONS

Co-administration of terfenadine, astemizole, or cisopride with LUYOX Tablets is contraindicated (see W LUYOX Tablets are contraindicated in patients with a history of hypersensitivity to fluvoxamine maleate. ne, asternizole, or cisapride with LUVOX Tablets is contraindicated (see WARNINGS and PRECAUTIONS)

WARNINGS
In petients receiving another serotoals respitate inhibitor drug in combination with monoamine oxidase inhibitors (MAOIs), there have been reports of serious, sometimes futal, reactions. 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. In addition, after stopping IUVOX° Tablets, at least 2 weeks should be allowed before setting a MAOI.
Terfonadine, astemizate and disciplide are all metabolized by the cytochrome P450IIIA4 isoenzyme. Increased plasmic concentrations of terfonadine, astemizate and disciplide case off prolongation and have been associated with torsades de points-type ventricular tedycurdia, sometimes futal. Although it has not been definitively demonstrated with flivoxomine is a petent IIIA4 inhibitor, it is likely to be. Consequently, it is recommended that flivoxomine not be used in combination with alther stemanties, associated and combination with alther stemanties, associated as of combination with alther stemanties.

polatis-type ventricular tachycardia, sometimes fatal. Although it has not been definitively demonstrated that flavoxamine is a potent IIIAA inhibitor, it is flavoy to be. Consequently, it is recommended that flavoxamine not be used in combination with either terbenodine, estematoly, or disperde.

Other Potentially important Drug Interactions
(Ass see PECALIMIDS - Dug Interactions) Exacedizacepines: Benzodizacepines metabolized by hepatic axidotion (e.g., alprazolam, midazolam, triazolam, etc.) should be used with caution because the clearance of these drugs is likely to be reduced by flavoxamine. Alprazolam-When flavoxamine metabolized by glouronicition (e.g., brazolam, conceptin, terracepoin) is unlikely to be diffected by flavoxamine. Alprazolam-When flavoxamine metabolized by application (e.g., brazolam ever exponomately hister has observed when riprazolam was administered alone; and placemate venerable placematers and application of application of applications are selected by about 50%. The elevated placem adequated placema adequated placematerians resulted in decreased psychomotop performance and memory. This interaction, which has not investigated using higher doses of flavoxamine, may be more pronounced if a 500 mg daily dose is condiministered of porticularly since flavoxamine exhibit non-interaction of LUVOX fables, the initial approach generally not observed by the designation of the process of the condiministered of plavoxamine exhibit non-interaction of LUVOX fables in the initial approach generally not observed places from a study in which healthy volunteers faxing 150 mg/dps and that of Hessenthyldizacepom to a level flavox more to low to measure over the course of the 2 week long study, it is likely that this experience significant that of subministered of the placematic of diazepom was not existent by 65% and that of Hessenthyldizacepom to level that was too low to measure over the course of the 2 week long study, it is likely that this experience significant of subministered of the placematic

PRECAUTIONS
General Activation of Manie / Hypomania: During premarketing studies involving primarily depressed potients, hypomania or mania occurred in approximately 1% of potients treated with througamine. Activation of mania / hypomania has does been reported in a small proportion of potients with motor affects desirated with a most activation of mania / hypomania has been reported in a small proportion of potients with a history of mania. Seizures: During premarketing studies, seizures were reported in 0.2% of throusamine-treated potients. LUVOX Tablests should be used continuely in primary depression or in association with monther primary disease who studies of the primary depression or in association with monther primary disease who is CD. Class supervision of high risk potients should accompany initial drug therapy. Prescriptions for LUVOX Tablests should be written for the smallest quantities to be a continuely and the primary depression or in association with writth Concomitation Hillings: Classify monitored clinical experience with LUVOX Tables in potients with decorate with control to tables to device the primary depression or the primary depression of the primary depression or device in Primary with Concomitation Hillings: Classify monitored clinical experience with LUVOX Tables in potients with decorate or conditions that could affect hemodynamic responses or metabolism. LUVOX Tables have not been evaluated or use products permarketing accorded from many clinical studies during the product's permarketing testing. Evaluation of the electrocardograms for potients with depression or OLO who porticipated in premarketing studies evalued to differences between flowarmountme and placebox in the mempersor of clinically importation, flowarmountme for Particular for Particular in potients with fiver before the initiation of treatment.

# Information for Patients

Information for Patients

Physicians are advised to discuss the following issues with patients for whom they prescribe LIVOX Tablets: Interference with Cognitive or Motor

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

Pergenancy: Patients should be obtained to notify their physicians if they become pregnant or intend to become pregnant during therapy with LIVOX Tablets.

Narsing: Potents receiving LIVOX Tablets should be odvised to notify their physicians if they are buest feeding on infant. See PECALITIONS - Nursing

Rothers.) Coeconitation: Medicarlone: Patients should be advised to notify their physicians if they are busing, or plan to take, any prescription or overtherecounter drugs, since there is a potential for clinically important interactions with LIVOX Tablets. Activates, with other psychotian redictions, patients should be advised to avoid alcohol while taking LIVOX Tablets. Allegeic Reactives: Patients should be advised to notify their physicians if they develop or onst, hives, or a related allergic phenomenon during therapy with LIVOX Tablets.

# Laboratory Tests

There are no specific laboratory tests recommended

# Drug Interactions

There one in specific idoptority tests recommended.

Drug Interactions:

There have been one postmarketing reports describing patients with weakness, hyperreflexio, and incoordination following the use of a selective serotonin nepitote inhibitor (SSR) and summittion. It concentrant heatment with summittation and an SSR (e.g., fluxetime, huvecomine, porceiving, settentials, critically warranteed, appropriate observation of the potential index official interactions with drangs that inhibitor or methods by Cytochrame P450 Inserpments Educed on a finding of substantial interactions of fluxocomine with certain drugs and limited in vitro date for the interactions of the recommendations and interaction in the involved in the metabolism of drugs such as warrant, heeplyfiline, certain benzodizegines and phenytoin. If UNION\* Tolles are to be administered together with a drug that is eliminated via outdative metabolism and has a narrow fluxogenies and phenytoin. If UNION\* Tolles are to be administered together with a drug that is eliminated via outdative metabolism and has a narrow fluxogenies and phenytoin. If UNION\* Tolles are to be administered together with a drug that is eliminated via outdative metabolism and has a narrow fluxogenies and phenytoin. If UNION\* Tolles are to be administered together with a drug that is eliminated via outdative metabolism and has a narrow fluxogenies and phenytoin. If UNION\* Tolles are to be administered together with a drug that is eliminated via outdative metabolism and has a narrow fluxogenies. In the continuous are recorded. Please see complete prescribing information to recommendations regarding UKS drugs such as monoromine administration of the properties of the clint and such that is eliminated via outdative metabolism and the properties of the clint and such as a fluxogenies of the clint

Programmy

First Speak: Effects - Programmy Category C: In terrology studies in rats and rabbits, daily and doses of fluvoxamine moleate of up to 80 and 40 mg/kg, respectively (approximately 2 lines the maximum human daily dose on a mg/m² basis) caused no fetal mailformations. However, in other reproduction studies in which pregnant rats were dosed through wearing there was 10 in an increase in pure martinity or both for separation of the control program of the potential benefit justifies the potential first to the fetuse.

Labor and Delivery

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

# Norsing Methers

As for many other drugs, flavoxamine is secreted in human breast milk. The decision of whether to discontinue unsing or to discontinue the drug should take into account the potential for serious adverse effects from exposure to fluvoxamine in the nursing infant as well as the potential benefits of LUVOX\* (howcomine enderle) blobbs thereby to the mother.

The efficacy of flavoxomine molecute for the treatment of Obsessive Compulsive Disorder was demonstrated in a 10-week multicenter placebo 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 fluvoxomine (see ADVERSE REACTIONS).

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

Generative Use
Approximately 230 portients participating in controlled premarketing studies with LUVOX Tablets were 65 years of age or over. No overall differences in safety were observed between these potients and varunger potients. Other reported clinical experience has not identified differences in response between the elderly and varunger potients. However, the clearance of fluoroximative is decreased by about 50% in whethy compared to younger potients potients. See Pharmacoliticities under CLINICAL PUBLINACOLOGIST), and genetic sensitivity of some older individuals also cannot be ruled out. Consequently, LUVOX tablets should be slowly tritted during initiation

# of therapy. ADVERSE REACTIONS

Associated with Discontinuation of Treatment
Of the 1087 OCD and depressed patients treated with fluvoxamine molecule in controlled clinical trials conducted in North America, 22% discontinued

tor the 1097 ULO and depressed primerits reteried with Individuals in controlled clinical mass controlled in north America, 22% asconsitudes tendented the tor on othersee event.

Adverse events in OCD Pedictric Population
In pedictric potients (N=57) heated with LUOX<sup>28</sup> Toldels, the overall profile of odverse events is similar to that seen in adult studies. Other reactions which have been reported in two or more of the pedictric potients, and were more frequent than in the placebo group (N=63) were characteristic more been reported in two or more of the pedictric potients, and were more frequent than in the placebo group (N=63) were characteristic more provided in the incidence in fundamental processors, provided in the incidence of 100 (Time-20) of the processors of the incidence of 100 (Time-20) of the processors of the incidence of 100 (Time-20) of the incidence, prosonnic, previousness, themse, processor, provided in controlled clinical processors, provided in the incidence of 100 (Time-20) of the incidence of 100 (Time-20) of the incidence, prosonnic, previousness, themse, processor, provided in the incidence of 100 (Time-20) of the incidence of 100 (T events were classified using a standard COSIARF-based Dictionary terminology. The prescriber should be aware that these flaures commot week to predict the incidence of side effects in the course of usual medical practice where patient characteristics and other fractors may differ from those that prevailed in the clinical thials. Similarly, the ched frequencies cannot be compared with figures obtained from other clinical investigations are nowlving different treatment, uses, and investigations the intelligence, however, to 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 the population studied. Adverses Events in OCD Placebo Controlled Studies While are Marked by Different (defined as at least a two-fold difference). In Rate from the Pooled Event Rates in OCD and obspression. Placebo Controlled Studies: The events in OCD studies with a two-fold difference in compared to event orders in OCD and obspression. Placebo Controlled Studies: While the very compared to event orders in OCD and obspression in COD studies with a two-fold increase in rate compared to event orders in OCD and obspression in COD studies with a two-fold increase in rate compared to event orders in OCD and obspression in COD studies with a two-fold increase in rate compared to event orders in OCD and obspression in COD studies with a two-fold increase in rate compared to event orders in OCD and obspression in COD studies with a two-fold increase in rate compared to event orders in OCD studies with a two-fold increase in rate compared to event orders in OCD studies with a two-fold increase in rate compared to event orders in OCD studies with a two-fold increase in rate compared to event orders in OCD studies with a two-fold increase in rate compared to event orders in OCD studies with a two-fold increase in the COD may observe the studies of the order of event orders in the COD may observe the studies of the order of event orders in the COD

Comparisons of fluvoramine molecute 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 portions meeting criteria for potentially important changes from baseline on various vital signs variables. The reversely not important differences between fluvoramine melanists and places.

revedled no important direteriors between inviscualities insulation of the Chaboratory Changes
Congarisons of fluvoximities mulecate and placebo groups in separate pools of short-term OCD and depression briols on (1) median change from baseline on various serum chemistry, hemotology, and urinalysis variables and on (2) incidence of patients meeting criteria for potentially important changes from baseline on various serum chemistry, hemotology, and urinalysis variables revealed no important differences between fluvoximities malerate and placebo.

ECG Changes

Comparisons of fluvoxamine molecute 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 gotients meeting criteria for potentially important changes from baseline on various ECG variables revealed no important differences between fluvoxamine molecute and placebo.

e 2: TREATMENT-EMERGENT ADVERSE EVENT INCIDENCE RATES BY BODY SYSTEM IN OCD AND DEPRESSION POPULATIONS COMBINED! (fluvoxamine [n=992] vs. placebo [n=778] by patients—percentage]: BODY AS WHOLE: Headache (22 vs. 20); Asthenia (14 vs. 6); Flu Syndrome (3 vs. 2); Chills (2 vs. 1). CARDIOVASCULAR: Polyintrions (3 vs. 2), DIGESTIVE SYSTEM: Nousea (40 vs. 14);

Tolkle 2: TREATMENT-HARGERIT ADVERSE EVENT INCIDENCE RATES BY BODY SYSTEM IN OCD AND DEPRESSION POPULATIONS COMBINED! (Incommine In-6972) is, ploade) (in-778) by printerin 5; pl. By Jordenia (B. s. 2); by Syndrom (B. s. 2); bills (Ver. 1). CARDIOVASCUIAR: Poliphorian (S. v. 2); bills (Ver. 1); Depression (10 vs. 3); Depospia (10 vs. 3); Anoexia (6 vs. 2); vonting (5 vs. 2); Floratere (4 vs. 3); Exotin (6 vs. 1); Anolyte (5 vs. 3); Properties (10 vs. 3); Properties (10 vs. 3); Properties (10 vs. 1); Properties (11 vs. 3); Properties (12 vs. 1); Properties (12 vs. 1); Properties (12 vs. 1); Properties (13 vs. 1); Properties (14 vs. 1); P

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

"Bosed on the number of tendles, "Bosed on the number of males,

Non-US Pastmarketing Reports

Voluntary reports of adverse events in patients taking LUVOX Tablets that have been received since tranket introduction and are of unknown causal relationship to LUVOX Tablets use include: toxic epidermal necrohysis, Stevens-Johnson syndrome, Henoch-Schoenlein purpura, bullous eruption, priopism, agranulocytosis, neuropathy, aplastic anemia, aneniy, acreation, hyponathemia, acute renal failure, hepatitis, and severe akinesia with fever when fluvoramine was co-dufmistrated with artitysycholic medication.

CAUTION: Federal law prohibits dispersing without prescription.

8E1252 Rev 3/97

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

# Pharmacia & Upjohn

Solvay **Pharmaceuticals** 

© 1998 Solvay Pharmaceuticals. Inc. All rights reserved.

SVL343

USI8453.00

January 1998

# EFFECTIVE FIRST-LINE SSRI THERAPY FOR OCD...



# EMERGING FROM THE PROFOUND ANXIETY OF OCD

# Low incidence of agitation

• 2% vs 1% for placebo<sup>1</sup>

# Low incidence of sexual dysfunction<sup>1</sup>

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

# **Favorable tolerability profile**

- Relatively low incidence of anticholinergic side effects in controlled trials of OCD and depression. LUVOX® Tablets vs placebo: dizziness 11% vs 6%; constipation 10% vs 8%; dry mouth 14% vs 10%¹
- 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%<sup>1</sup>
- 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 recommended¹

\*Parameters occurring ≥ 1% with fluvoxamine maleate.

Please see brief summary of prescribing information on adjacent page.



**AVAILABLE IN 25-mg TABLETS** 

First-line S.S.R. Inite henrageny for obsessions and compulsions