## **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

LUVOX Tablets are indicated for the treatment of obsessions and compulsions in patients with Obsessive Compulsive Disorder (OCD), as defined in the DSM-HIFA. 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, astemizale, or cisapride with LUVOX Tablets is contraindicated (see WARNINGS and PRECAUTIONS). LUVOX Tablets are contraindicated in patients with a history of hypersensitivity to fluvoxamine maleate.

### WARNINGS

WARNINGS
In patients receiving another serotonin reuptake inhibitor drug in combination with monoamine oxidase inhibitors (MAOIs), there have been reports of serious, sometimes fatal, 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 LUYOX® Tablets, at least 2 weeks should be allowed before starting a MAOI.

Tarfanadine, estemizade and disapride are all metabolized by the cytodrome P450IIIA4 isoenzyme. Increased plasma concentrations of terfanadine, astemizade and disapride cause QT prolongation and have been associated with torsades beolists-type ventricular tadycardia, sometimes fatal. Although it has not been definitively demonstrated that fluvoxamine is a potent IIIAA inhibitor, it is likely to be. Consequently, it is recommended that fluvoxamine not be used in combination with either terfanodine, astemizade, or cisapride.

Other Patentially important Drug Interactions

(Also see PRECAUTIONS - Drug Interactions) Benzadiazepines: metabolized by phopic oxidation (e.g., alprazolam, midazolam, nutrazolam, etc.) Sound be used with cutom because the clearance of hese drugs is likely to be reduced by fluvoxamine. The Genance of benzadiazepines metabolized by glucuronidation (e.g., lorazopam, corzepom, tenrazepom) is unlikely to be effected by fluvoxamine. Alprazolam-When fluvoxamine meleste (100 mg gd) and alprazolam produces and the phomacokinetic parameters (AUC, Ca., I.) of alprazolam was optimisted and cleanance was reduced by doors OSis. The

metabolized (10) ang do and quizodinal (1 mg qil) were condiministered in seculty strie, plasma concentrations and other paramosine-white involvations (1 mg qil) were condiministered in seculty strie, plasma concentrations and other paramosines (AUC, Co., I.) of alprazolam were approximately hive those observed when alprazolam was administered alone; and cleanace was reduced by about 50%. The elevated plasma alprazolam, and the plasma objects of the second mission of the plasma objects of the second plasma of the second plasma objects of the second plasma objects of the second plasma of the second plasma objects of the second plasma of the second plasma objects of the second plasm

General Activation of Mania/Hypomania: During premarketing studies involving primarily depressed patients, hypomania or manio occurred in approximately 1½ of patients treated with fluoroxamine. Activation of mania/hypomania has also been reported in a small proportion of patients with major affective disorder who were treated with other marketed antidepressorst. As with all antidepressors, LIVOX for fluoroxamine-treated patients. LIVOX flowing because the second of a 2½ of fluoroxamine-treated patients. LIVOX flowing account of a country of the possibility of a suicide attempt of the patients with a history of seizures. It should be discontinued in any patient who develops seizures. Suicide: The possibility of a suicide attempt. cautiously in potients with a history of seizures. It should be discontinued in any potient who develops seizures. Suicide: The possibility of a suicide attempt is inherent in potients with depressive symptoms, whether hese occur in primary depression or in association with another primary disorder such as CO. Close supervision of high nisk potients should occurrency within drug therapy. Prescriptors for LIVOX Tables should be written for the smallest quantity of totalest consistent with good patient management in order to reduce the risk of overdose. Use in Parlients with Concomitant Illness; Closely monitored clinical experience with LIVOX Tables in potients with concomitant systemic illness is intented. Caution is advised in administering LIVOX Tables to patients with diseases or conditions that could offer hemodynamic responses or metabolism. LIVOX Tables have not been evaluated or used to any appreciable extent in potients with a recent history of myocardial inforction or unstable heard disease. Potients with these diagnoses were systematically excluded from many clinical studies during the product speriorecking is testing. Foundation of the electrocardiagnorms for patients with depression or OD who participated in permatchering studies revealed no differences between fluvoxornine and placebo in the emergence of clinically important ECG changes. In patients with liver dysfunction, fluvoxornine clearance was decreased by approximately 30%. LIVOX Tablest should be slowly thrated in patients with liver defunction during the initiation of trustment. 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

Parformance: Since any psychoctive drug may impair judgement, thinking, or motor skills, patients should be coultoned 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 advised to notify their physicians if they accome pergenator in tended to become pregnant during therapy in LUVOX Toblets.

Narsing: Potients receiving LUVOX Toblets should be advised to notify their physicians if they are breast feeding an infant. (See PREAUTIONS - Nursing

Mothers). Concomitant Medication: Patients should be advised to notify their physicians if they are bicking, or join to toke, any prescription or overtherounter drugs, since there is a potential for clinically important intercolors with LUVOX Toblets. Acknob. As with other psychiatric medications,
patients should be advised to avoid alcohal while taking LUVOX Toblets.

Allergic Reactions: Patients should be advised to notify their physicians if they develop a rash, hives, or a related allergic phenomenon during therapy with LUVOX Toblets.

Laboratory Tests
There are no specific laboratory tests recommended.

Drug Interactions
There have been true postmakeling reports describing patients with weakness, hyperrellexia, and incoordination following the use of a selective serotomic response in the patient of the patient is advised. Patential interactions with drugs that inhibit or are Metabolized by Cytochrame P450 Isozymes: Based on a Indiag of substantial interactions with drugs that inhibit or are Metabolized by Cytochrame P450 Isozymes: Based on a Indiag of substantial interactions with drugs that inhibit or are Metabolized by Cytochrame P450 Isozymes: Based on a Indiag of substantial interactions with drugs that inhibit or are Metabolized by Cytochrame P450 Isozymes: Based on a Indiag of substantial interactions with a metabolism of laws; such as variorial, theophyline, certain peacepast that invocarione inhibits isochrames that or known to be involved in the metabolism of laws; such as variorial, theophyline, certain peacepast; metabolism of laws; and a programoid. A clinically significant flavocarione interaction is possible with drugs having a narrow therapeutic ratio such as tertained in a continuous cont

Cardinogenesis. Muragenesis, Impairment of Fertility and operation of fertility with fluvoxomine maleate, There was no evidence of corinogenicity, mutagenicity or impairment of fertility with fluvoxomine maleate for 20 (females) or 26 (males) norths. The dairy does in the high does groups in these studies were increased over the course of the study from a minimum of 160 mg/kg to consistent. The dairy does in the high does groups in these studies were increased over the course of the study from a minimum of 160 mg/kg to consistent of 240 mg/kg in chapters. The maximum of 240 mg/kg is oppoximately 5 times the maximum human daily dose on a mg/m² basis. Mutagenesis: No evidence of mutagenic potential was observed in a mouse microardices test, on it with ord formersome observation and female rise, up to 160 mg/kg in chapters or without metabolic activation. Impairment of Fertility: In lettility studies of male and female rise, up to 80 mg/kg and group or will be fluvoxomine maleate, (approximately 2 times the maximum human daily dose on a mg/m² basis) had no effect on mating performance, duration of gestution, or pregnancy rate.

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

Pregnancy

Teratogranic Effects - Pregnancy Category C: In teratology studies in rots and rabbits, daily and doses of fluvoxamine maleate of up to 80 and 40 mg/kg, respectively (approximately 2 times the maximum human daily dose on a mg/m² basis) caused no fetal malformations. However, in other productions addes in which pregnant are was consumed as was (1) an increase in purp motality at this flosen at 30 g/kg and above but not at 20 mg/kg), and (2) decreases in postnatal pup weights (seen at 160 but not at 80 mg/kg) and survival (seen at all doses; lowest dose tested = 5 mg/kg). (Doses of 5, 20, 80, and 160 mg/kg are approximately 0, 1, 0.5, 2, and 4 himse the maximum human daily dose on a mg/m² basis.) While the results of a crossfortaming sulty implied that the test same of these sensit likely counted secondarily not material toxicity on dote of adirect day effect on the fetures or pups could not be ruled out. These are no adequate and well-canholled studies in pregnant women. Fluvoxamine maleate should be used during pregnancy only if the potential benefit justifies the potential task to the feture.

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

## Nursing Mothers

NOTIFIED MONITORS.

As for many other drugs, fluvoxamine 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 fluvoxamine in the nursing infant as well as the potential benefits of LUYOX\* (fluvoxamine maleate) Tablets therapy to the mother.

### Pediatric Use

Pediatric Use
The efficacy of fluvoramine maleate 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 odult studies with fluvoramine (see AUVENSE REACTIONS).

Decreased appetric and weight last stave been observed in association with the use of fluvoramine 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

Genariar Use
Appointmet 9, 230 patients participating in controlled premarketing studies with LUVOX Tablets were 65 years of age or over. No overall differences in safety were observed between these patients and younger pointers. Other reported clinical experience has not identified differences in response between the eldorly and younger pointers. Other reported clinical experience has not identified differences in response between the eldorly and younger patients. However, this cleanance of floroxomine is decreased by about 50% in elderly compared to younger patients (see Pharmacolimetrs under CUNICAL PHARMACOLOT), and greater sensitivity of some older individuals also cannot be ruled out. Consequently, LUVOX habitest should be Sowly thatted during initiation.

### ADVERSE REACTIONS

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

Adverse events in CVD Pediatric Population In pediatric potents (IH=57) teated with LUNOVE fablets, 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 pediatric patients, and were more frequent than in the placebo group (IN=63) were: abnormal thinking, cough increase, dysmenorihea, ecclymosis, emotional lability, existica's, hyperkinesio, affection, manic reaction, restly, sinusitis, and weight develose.

increase, dysmenormen, echymoxis, emofared lability, epistaxis, Pyperkinesio, infection, mainic reloction, risth, sinustits, and weight decrease. Events for which the incidence in fluoroamine malerate was equal to or less than the incidence in placebo (Ni-63) and involved two or more of the pediatric study patients were: abdominal pain, abnormal dreams, lever, headache, nouses, nervousness, pain, pharrygitis and hinitis.

Incidence in Controlled Trials - Commonly Observed Adverse Events in Controlled Clinical Trials: LIVIVIX foldets have been studied in controlled this of 000 (19–320) and depression (ni-1350). In general, observe event trate were similar in the two datos set. The most commonly observed adverse events associated with the use of LIVIVIX foldets and kikely to be drug-related (incidence of 5% or greater and or less than the placebod power form Table 2 were: sommolence, insomain, nervousness, hemor, nauses, dyspepsia, anorexie, vomiting, abnormal ejeculation, astheria, and sweding in a good of two studies involving only polients with OCI, the following additional events were identified using the above rule: dy mount, decreased libido, unnour frequency of 1% or integrancy of 1% or moe, and were more frequent than in the placebod gover, among patients treated with LIVIVIX foldets in two short term piecebo cannolled ODI table (10 week) and depression trials (6 week) in which polients 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 notego. In the internet integration of the treatment, Reported adverse events were classified using a standard COSIRR\*hoseb Dictionary terminalogy. The prescribe should be owner than three figures cannot be used to predict the incidence of side effects in the cause of usual medical practice where patient characteristics and other factors may differ internment. Reported adverse events were classified using a standard COSIRR\*hoseb Dictionary terminalogy. The prescriber should be owner than three f in a future of use entails. In the custor we to soon mentury pounds where purious multicularities and understance and understance that the futured intide. Similarly, the crite of frequencies comoto be compared with figures obtained from other clinical investigations involving different neutrements, uses, and investigations. The critical figures, however, do provide the prescribing physicion with some bosts for estimating the relative contribution of drug and non-drug factors to the side-effect incidence rate in the population studied. Adverse Events in OCD Placebo Centralled Studies Which are Markedly Different (defined as at least a two-fold difference) in Rate from the Pooled Event Rates in OCD and Depression Placebo Controlled Studies: The events in OCD studies with a two-fold decrease in our compact event rather in OCD and population of Placebo Controlled Studies: The events in OCD studies with a two-fold decrease in our compact or event rather in OCD on depression studies were obtained, obnormed ejaculation (mostly blurred vision), Additionally, there was an approximate 25% decrease in nousea. The events in OCD studies with a two-fold inclease in note compared to event rather in OCD and depression studies were, ostheria, obnormed ejaculation (mostly delayed ejaculation), avaiety, infection, thinitis, amagazini of modes), depression, bladd decreased, phanyrapis, agritation, impotence, myodonus/hvitit, thirst, weight loss, leg camps, myodon and univary relention. These events are listed in order of decreased greater in OCD thirds.

Vital Sign Changes

Companying of the hovergraine melecte and elevative grows in secrette mode of short-term PCD and depression trick on (1) melion change from boseline on

Comparisons of Bhoxamine maleate and placebo groups in separate pools of short-term DCD 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.

Laboratory Changes
Comparisons of flowcomine molecule and placebo
Comparisons of flowcomine molecule and placebo groups in separate pools of short-term OCD and depression trials 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 flowcommine malaote and placebo.

Comparisors of flavoxomine maleate and placebo groups in separate pools of short-term OCD and degression trials on (1) mean change from baseline on various ECG variables and on (2) incidence of patients meeting criteria for patentially important changes from baseline on various ECG variables revealed na impartant differences between fluvoxamine malente and placeho

TREATMENT-EMERGENT ADVERSE EVENT INCIDENCE RATES BY BODY SYSTEM IN OCD AND DEPRESSION POPULATIONS COMBINED (fluvoxumine [n=892] vs. plocebo [n=778] by potients—percentage): BODY AS WHOLE: Headache (22 vs. 20). Asthenia (14 vs. 6); Flu Syndrome (3 vs. 2); Chills (2 vs. 1). CARDIOVASCULAR: Polpitutions (3 vs. 2). DIGESTIVE SYSTEM: Nausea (40 vs. 14). Actheria (14 vs. 6); Hu Syndrome (3 vs. 2); Chillis (2 vs. 1). CARDIOVASCULAR: "Foliatrions (3 vs. 2). DOIESTIVE SYSTEM: Nausse (40 vs. 14). Diarrher (11 vs. 7); Contapion (10 vs. 8); Dyspepsio (10 vs. 5); Anoreoi (6 vs. 2); Volming (5 vs. 2); Filatriog (5 vs. 2); Pilatriog (5 vs. 1); Pilatriog (5 vs. 3); Pilatriog (5 vs. 1); Pilatriog (7 vs. 3); SPECIAL SENSES: Totale Prevession (3 vs. 1); Amblyopin (3 vs. 2); Impotence (2 vs. 1); Amblyopin (3 vs. 2); Impotence (2 vs. 1); Pilatriog (2 vs. 1); Pilatriog (2 vs. 1); Pilatriog (2 vs. 1); Pilatriog (5 vs. 2); Pilatriog (5 vs. 2); Pilatriog (5 vs. 1); Pilatriog (5 vs. 2); Pilatriog (5

(7 vs. 3). SPECIAL SENSES: lasts Pervession (3 vs. 1): Amblyopin (3 vs. 2): LINGERINITAL: Abnormal Epoculonian (8 vs. 1): Ultimary Frequency (3 vs. 2): Imported (2 vs. 1): Amblyopin (2 vs. 0): Ultimary Retention (1 vs. 0):

Events for which fluoxomine moderal encidence was equal to or less than placebo are not listed in the table above, but include the following: addominal point, denormal dearns, appetite increases, back pain, cheer point, ordersion, dysmenorthee, fever, infection, leg carrays, magnine, mydigin, pain, paesthesis, phorygating, sociated hypotension, prinsip, sach, hindis, hists and hinnish. Ricubes "Tombords," "Tombet arcticion and abscass," and "creas." Mustry feeling warm, hot, or flushed. "Mostly "deleyed ejeculation." "Includes "Tombords," "Tombet arcticion and abscass," and "creas." Mustry feeling warm, hot, or flushed. "Mostly "deleyed ejeculation." "Includes "Tombords," "Tombet arcticion and abscass," and "creas." "Mustry feeling warm, hot, or flushed. "Mostly "deleyed ejeculation." "Includes a common and abscass," and "creas." "Mustry feeling warm, hot, or flushed. "Mostly "Abscass" and a fluoridation of a 1973 political exposures in polients suffering COL or Mujor Depressive Disorder. Ultroward events associated with filis exposure were recorded by furition pressive disorder. Includes a common and the late political provides or mensingful estimate of the proportion of individuals exposures in political sufficiency of the provides or mensingful estimate of the proportion of the 2737 political responsibility of the proportion of the 2737 political exposures to multiple doses of fluovocamine molecule were severed and provided and provided and recommon and provided and pro

Based on the number of females, Based on the number of males,

### **Non-US Postmarketing Reports**

New-up-rostmankening keeports Voluntary reports of oderese events in patients taking LUVOX Tablets that have been received since market introduction and are of unknown cousal relationship to LUVOX Tablets use include: toxic epidermal necodysis, Stevens-Johnson syndrome, Henoch-Schoenlein purpura, bullous eruption, priapsim, organilus/pross, neuropathy, oplastic anemia, anaphylocitic reaction, hyporatientia, ccute renal failure, hepatitis, and severe ckinesia with fever when flavoxamine was coordimistered with antipsychotic readication.
CAUTIONE: Federal law prohibits dispensing without prescription.
8F1952 Res 3/95.

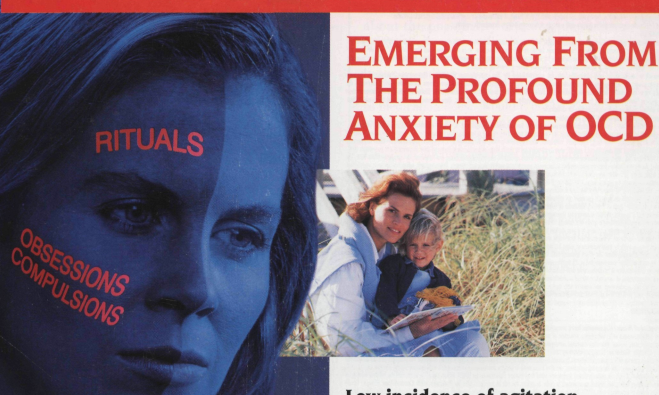
Reference: 1. Data on file, Solvay Pharmaceuticals, Inc.

## Pharmacia&Upjohn

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





• 2% vs 1% for placebo

# 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%1
- 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



**AVAILABLE IN 25-mg TABLETS** 



\*Parameters occurring ≥ 1% with fluvoxamine maleate. Please see brief summary of prescribing information on adjacent page.