## LUVOX <sup>40</sup> (fluvaxamine 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

HIDSCATTOR'S AND GONOL LUVOX tables are indicated for the treatment of absessions and compulsions in patients with Obsessive Compulsive Disorder (OCD), as defined in the DSMHIP. Obsessive Compulsive Disorder is characterized by recurrent and persistent ideas, thoughts, impulses or images (absessions) that are ego-dystoric and/or repetitive, purposeful, and intentional behaviors (compulsions) that are recognized by the person as excessive or unreasonable. and/or repetitive, purposetu CONTRAINDICATIONS

Condiministration of terfenodine, astemizale, or cisopride with LUVOX Tablets is contraindicated (see WARNINGS and PRECAUTIONS). LUVOX Tablets are contraindicated in patients with a history of hypersensitivity to fluvoxamine malecte.

## WARNINGS

WARNINGS In patients receiving another serotonin reuptake inhibitor drug in combination with monoamine oxidase inhibitors (MAOIs), there have been reports of serious, sometimes total, 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 LUVOX® Tablets, at least 2 weeks should be allowed before starting a MAOI. Terfenadine, astemizole and disapride are all metabolized by the cytachrome P450111A4 isoenzyme. Increased plasma concentrations of terfenadine, astemizole and cisapride cause QT prolongation and have been associated with torsades de points-type ventricate tachycordia, sometimes fatal. Althoogh thas not been definitively demonstrated that fluvoxamine is a potent IIA4 inhibitor, it is likely to be. Consequently, it is recommended that fluvoxamine not be used in combination with either terfenadine, astemizoles or cisapride.

points-type ventricular tachycardia, sometimes tatia, Although if has not been deminitively demonstrated material transmission is a potent likely his likely to be consequently, it is recommended that fluvoxamine not be used in combination with either terfendines, astemizole, or dispride. Other Potentially Insportant Drug Interactions (Also see REGAM), it is recommended that fluvoxamine not be used in combination with either terfendines, astemizole, or dispride. Other Potentially Insportant Drug Interactions (Also see REGAM), it is recommended that fluvoxamine. The cleance of bearcolour, priors and the potential of the cleance of these drugs is likely to be relaxed by throwamine. The cleance of bearcolourspines methodically to be predived by throwamine. The cleance of bearcolourspines method (100 mg d) and dpurzalam (11 mg did) were codministered to stady state, plasma concentrations and bree pharmocknets purchased processes and updurzalam was administered alone; ond cleance was reduced by class 15%. The elevated pharma dpurzalam cleance those deserved when dpurzalam was administered alone; ond cleance was reduced by class 15%. The elevated pharma dpurzalam cleance that and a state of the second state of t

## General

Activation of Mania/Hypomania: During premarketing studies involving primarily depressed patients, hypomania or mania occurred in approximately /// or patients treated with fulveromine. Activation of mania/hypomania has also been reported in a small proportion of patients with major affective disarder who were treated with there marketed antidepressment. Sk with all antidepressments, LUVOX tablets should be used cardinary in patients with a cardinary of saizues. It should be discontinued in any proximately is inherent in patients with depressive symptoms, whether these accur in primary depression or in association with another primary disorders such a Suckider. The possibility of a suicide attempt is inherent in patients with depressive symptoms, whether these accur in primary depression or in association with another primary disorders such as SUCION Tablets should be written for the smallest quantity of tablest consistent with also of patient management in order to reduce the risk of overdorse. Use in *Patientess with Concomitant Hinesse*: Casely monitored clinical experience with LUVOX tablets in patients with concomitant systemic illness is limited. Caution is advised in odministening LUVOX Tablets to patients with descess or conditions ther could affer therrodynamic responses or metabolism. LUVOX tablets have not been avoluted or used to any appreciable extent in patients with a creath story of myocardial infrarction or unstable herd disease. Pranets with degressions were systematically patients with these dispatients during there are also parametering testing. Luvolation of the democardiograms or 4D0 with these outpatient or 4D0 with a systemic illowed or used to any appreciable extent in patients with a creath story of myocardial infrarction or unstable herd disease. Pranets with degression or 0D0 who patients with the dystanction, fluevacamine clearance was decreased by approximately 30%. LUVOX Tablets should be should be should be down'thanded in patients with liver dystanchand uning the ini dysfunction during the initiation of treatment. Information for Patients

Physicians are advised to discuss the following issues with patients for whom they prescribe LUVOX Tablets: Interference with Cognitive or Motor Physians are advised to discuss the following issues with patients for whom they prescribe LUVOX Tablets. Interference with Cognitive or Motor Performance: Since any psychoactive drug may impair judgement, thinking, or motor skills, patients should be acutioned about operating hazardous machinery, including automobiles, until they are certain that LUVOX Tablets theory does not downsky diffet their ability to engage in such activities. Pregnancy: Patients should be advised to notify their physicians if they become pregnant or intend to become pregnant during theory with LUVOX Tablets. Narsing: Patients ineaving LUVOX Tablets should be advised to notify their physicians if they are taking, or plan to take, any prescription or over the counter drugs, since there is a patiential for chincilly important interactions with LUVOX Tablets. Alcohoot: As with other psychotopic medications, potents should be advised to notify their physicians if they are taking, or plan to take, any prescription or over the counter drugs, since there is a patiential for chincilly important interactions with LUVOX Tablets. Alcohoot: As with other psychotopic medications, potents should be advised to rotify their physicians if they are taking, or plan to take, any prescription or over the counter drugs, since there is a patiential for chincilly important interactions with LUVOX Tablets. Machoot: A with other psychotopic medications, develop a rash, hives, or a related allergic phenomenon during therapy with LUVOX Tablets.

## Laboratory Tests

There are no specific laboratory tests recommended.

There are no specific laboratory tests recommended.
Drug Interactions
There have been are postmarketing reports describing patients with weakness, hypereflexia, and incoordination following the use of a selective sectorian
requise inhibitor (SSR) and sumatipitor. If concomiant treatment with sumatriptor and an SSRI (e.g., fluxostine, fluxos

studies stubilishing the benefits or risks of combined use of ECT and fluxoxamine molecte. Carcinogenesis, Motagenesis, Impairment of Fertility Carcinogenesis: There is no evidence of carcinogenicity, mutagenicity or impairment of fertility with fluxoxamine maleate. There was no evidence of carcinogenesis: There is no evidence of carcinogenicity, mutagenicity or impairment of fertility with fluxoxamine maleate. There was no evidence of carcinogenesis: There is no evidence of carcinogenicity, mutagenicity or impairment of fertility with fluxoxamine maleate. There was no evidence of (males) months. The daily 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 a maximum of 240 mg/kg in tast, and from a minimum of 135 mg/kg to a maximum of 240 mg/kg in hometers. The maximum does of 240 mg/kg is oppositoriately 6 times the maximum human daily does on a mg/m<sup>2</sup> bois. Mutagenesis: No evidence of mutagenic potentil does observed in a mouse incorructers test is on in vitro chromoseme abereficion test, or the Amse minicadi mutagen test with a without metabolic activation. Impairment of Fertility: In fartility studies of male and female rats, up to 80 mg/kg/dey orally of fluxoxamine maleate, (approximately 2 times the maximum human daily does on a mg/m<sup>2</sup> basis) had no effect on mating performance, duration of gestation, or pregnancy role. Prevenency

## Prognancy

Programy *Teratogenic Effects - Pregnancy Category C:* In teratology studies in rats and rabbits, daily and doses of Ruvoxamine maleate of up to 80 and 40 mg/kg, respectively (caparoximately 2 times the maximum human daily dose on a mg/m<sup>2</sup> basis) caused no fatal malformations. However, in other reproduction studies in which pregnant rats were dosed through verning there was (1) an increase in pup mortality at birth (seen at 80 mg/kg and but and 120 mg/kg), and (2) decreases in postantial puweighs (seen at 160 but not at 80 mg/kg) and suvivations tables in other = 5 mg/kg). (Doses 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/m<sup>2</sup> basis.) While the results of a cross-fostering study implied that at least same of these results likely occurred sacandarily to maternal toxicity, the role of a direct day effect on the therase or purs caudion to be red out. There are no dequare and well-controlled studies in pregnant women. Huroxamine maleate should be used during perganacy only if the potential benefit justifies the potential risk to the fetus. Labor and Delivery he effect of Huxonnine on labor and delivery in humans is unknown.

The effect of fluvoxamine on labor and delivery in humans is unknown

## **Nursing Mothers**

As for many other days, fluwaramine is secreted in human breast milk. The decision of whether to discontinue mursing or to discontinue the drug should take into account the potential for serious adverse effects from exposure to fluwaramine in the nursing infant as well as the potential benefits of LUVOX\* (fluvoxamine maleate) Tablets therapy to the mother.

## Pediatric Use

Treatment as a flux stantine maleate for the treatment of Obsessive Compulsive Disorder was demonstrated in a 10-week multicenter placebo astholled study with 120 autyatients ages 3-17. The adverse event profile observed in that study was generally similar to that observed in adult studies with flaxonamic (see AdVERSE REALTIONS).

Decreased appetite and weight loss have been abserved in association with the use of fluvoxamine 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.

### Gerintric Use

Genative Use Approximately 230 patients participating in controlled premarketing studies with LUVOX Tablets were 65 years of age or over. No overall differences in sofery were observed between these patients and younger portents. Other reported clinical experience has not identified differences in regiones between the elderly and younger patients. However, the clearance of throncommers is decreased by about 50% in elderly composed to younger patients (see Pharmacokinetics under CUNICUL PHARMACUOGE) and greater saretistivity of some older individual also camon be hald out. Consequently, LUVOX holbes shuld be shuld had uning initiation of therapy

## 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 OCD Pediatric Population to pediatric patients (N=57) treated with LUVOX® Tablets, the overall pratile of adverse events is similar to that seen in adult studies. Other reactions which In provide the partial in two or more of the pediatric patients, and were more frequent than in the placebo group (N=63) were: charmal thinking, cough increase, dysmenorhea, eachymasis, emotional lability, epistaxis, hyperkinesia, infection, manic reaction, rash, sinusitis, and weight decrease.

Indicate, gradientarina, exclusione encloses encloses and an enclose encloses and e odverse events associated with the use of LUVOX Tablets and likely to be drug-related (incidence of 5% or greater and at least twice that for placebo) derived from Table 2 were: commolence, incomnia, pervosaness, theman, nausen, dyspepsia, manexia, vannitrag, admarmal ejaculation, actimating, and sweating, in a pool of the subles involving only prientens with OCD. In following additunal events were identified using the above rule: dy mouth, decreased liable, unitary frequency, anargosmis, rhinitris and taste perversion. **Adverses Events Occurring et an incidence of** 15%; Table 2 anotes, and the submitted using the above rule: dy mouth, decreased liable, unitary frequency, anargosmis, rhinitris and taste perversion. **Adverses Events Occurring et an incidence of** 15%; Table 2 anotes, and the observation and an event and the placebo group, anong potients theaded with LUVOX Tablets in two short term placebo controlled OCD tricks (10 week) and depension tricks (6 week) in which potients were dosed in a rung of generally 100 to 300 mg/day. This table shows the percentage of primers in each group who had at least one occurrence of one event at one time during their theatment. Reported adverse events were clossified using a standard OSISMFbased Dictionary terminology. The prescube should be avore that these figures connot be used to predict the incidence of side effects in the course of used unadical practice where potient characteristics and other factors may differ from those that prevailed in the clinical triaks. Similarly, the cited frequencies cannot be compared with figures obtained from other clinical investigations involving ifferent theatments, uses, and investigators. The cited figures, however, do provide the prescripting physion with some basis for estimating the relative contribution of drug and nondrug factors to the sideeffect incidence rate in the population studied. Adverse Events in OCD Placebo Controlled Studies Whick are **Markedly Different (defined as st least a two-fold difference) in Kate from th** Marked of the series of the se

Introduction and provinces interest or end or a contract or accounting section in the CC matrix. Virtid Sign Changes Comparisons of fluwacamine malante and placebo groups in separate pools of short-term OLD and depression trials on (1) median change from baseline on various virtal signs variables and on (2) incidence of potentis meeting artients for potentially important changes from baseline on various virtal signs variables revealed on important differences between fluwacamine malante and placebo.

Laboratory Changes Comparisons of fluxoxamine molecte and placebo groups in separate pools of short+erm OCD and depression trials on (1) median change from baseline on various serum chemistry, hematology, and urinalysis variables and an (2) incidence of patients meeting criteria for potentially important changes from baseline on various serum chemistry, hematology, and urinalysis variables revealed no important differences between fluvoxamine malecte and placebo. ECG Changes

Comparisons of fluvaxamine maleate and placebo groups in separate pools of short+erm DCD and depression trials on (1) mean change from baseline on various ECG variables and no (2) incidence of patients meeting artiferia for potentially important changes from baseline on various ECG variables revealed no important affarences between fluvarrine marketen and placebo.

Notes Concernent (2) inclusion of the present of them to poletimate implorted transport of the section absent of the concernent of the present of the section of the present of the section of the present of the section of the section of the present of the section of the present of the section of the section of the present of the section of the sec

Other twents Observed During the Premarkering Evoluation et LUVOX tablets During premarkering clikic mits conducted in Noth America and Europe, multiel doess of Ilucoxomine moleate were administered for a cambined total d 2737 patient exposures in patients suffering OCD or Major Depressive Disorder. Unhowed 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 individuol experiencing adverse events without first grapping similar types of univoard events into a limited (i.e. geduced) number of standard event cragonies. In the tabulations which follow, a standard COSMR/Isosed Dictionary terminology has been used to dossify reported devents over an induced in the list below, with the following exceptions: 1) those events of ended ore occssion while receiving Duroxamine maleate. All reported events are included in the list below, with the following exceptions: 1) those events of ended y listed in table 2, which tabulates indexec trates of common adverse experiences in placet-bocramoted OL and depression clinical trates, are excluded. 2) those events for which a drug case was considered remote (i.e., neoplasia, gastrantestrind cartionen, heppes simplex, heppes zotet, opplication si the exotion, and using exclusion clinical diverse on using a center second in diverse positient and judget to not be potentially evidents on a less of 1/100 prients, inferent fuevoarnine moletue has not been established. Events as chritter dissilied within body system categories on denumerated in oder of decreasing frequency using the following definitors: frequent doverse reveits are defined as flows couring on one or none occssion is and less 1/100 prients. Body see with see response to the couring between 1/100 and 1/1000 prients; and readeness events are trate accurating in less tables. Julioo and the case seents are those accurating between 1/100 and 1/1000 prients; disexe kidnev calculus, hematospermia<sup>2</sup>, oliquria. Based on the number of females, "Based on the number of males

Based on the number or remotes: based on the numbers on numes. Non-US Postmarketing Reports Voluntary reports of otherse events in patients taking LUYOX Toblets that have been received since market inhoduction and are of unknown causal relationship to LUYOX Tables use include: taxic epidemial neculopis, Stevens-Johnson syndrome, Herach-Schoenlein purpura, ballous euption, prinjarm, organulacytosis, neuropathy, palacit, ameria, anaphylacitric reaction, hyponatienina, acute renal failure, hepotitis, and severe akinesia with lever when fluvoxamine was ro-administered with antipsychiat medication.

CAUTION: Federal law prohibits dispensing without prescription 8E1252 Rev 3/97

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



Solvay Pharmaceuticals

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# **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'

 LUVOX<sup>®</sup> 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<sup>1</sup>: dizziness 11% *vs* 6%; constipation 10% *vs* 8%; dry mouth 14% *vs* 10%
- 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>
- Concomitant use of LUVOX® Tablets and monoamine oxidase inhibitors is not recommended<sup>1</sup>

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

# First-line SSRI therapy for obsessions and compulsions

fluvoxamine maleat

25 mg TABLETS 50 mg & 100 mg SCORED TABLETS

RITUALS