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 Tables are indicated for the treatment of obsessions and compulsions in patients with Obsessive Compulsive Disorder (OCD), as defined in the DSM-III-R. 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.

Co-administration of terfenodine, astemizale, or cisagride with LUVOX Tablets is contraindicated (see WARNINGS and PRECAUTIONS) LUVOX Tablets are contraindicated in patients with a history of hypersensitivity to fluvoxamine malec

WARNINGS

Transmired receiving another serotonin reuptake inhibitor drug in combination with monoamine oxidase inhibitors (MAOIs), there have been reports of serious, sometimes fotal, reactions. Therefore, it is recommended that LUYOX* 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.

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Terfenadine, a scienziale and disparide are all metabolized by the cytodrome P450IIIA4 isoenzyme. Increased plasma concentrations of terfenadine, astemizale and disparide cause QT prolongation and have been associated with forstades points-type ventricular tackyradiic, sometimes fatol. 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, astemizale, or disparide.

Other Potentially Important Drug Interactions: Benzodiazepines: Benzodiazepines metabolized by hepotic oxidation (e.g., alprazolam, midazolam, rizarolam, etc.) should be used with counts because the leasurace of these drugs is likely to be reduced by fluvoxamine. The charance of benzodiazepines metabolized by glocuronidation (e.g., lorazepam, oxazepam, temazepam) is unlikely to be affected by fluvoxamine. The charance of benzodiazepines metabolized by glocuronidation (e.g., lorazepam, oxazepam, temazepam) is unlikely to be affected by fluvoxamine. Alprazolam—When fluvoxamine maletet (100 mg agd and diurzalam (1 mg qid) were condiministered to stody stute, plasma concentrations and other pharmacokinetic purameters. Alprazolam was offinized plasma alprazolam concentrations resulted in decreased psychomotor performance and memory. This interaction, which has not been investigated using higher doss of fluvoxamine, may be more pronounced if a 300 mg aduly dose is co-administered, aprincularly since fluvoxamine characterial and intration to the lowest effective dose is recommended. No dosage adjustment is required for LIVOX Tablets. Diazepam—The co-administration of LIVOX Tablets, the initial alprazolam dasage should be at least however, and alargapine is generally not advisoble. Because fluvoxamine redu administered. Theophylline: The effect of steady-state fluvoxamine (50 mg bid) on the pharmacokinetics of a single dose of theophylline (375 mg as 442 mg aminophylline) was decreased approximately 3-fold. Therefore, if theophylline is co-administered with fluvoxamine malente, its dose should be reduced to one third of the usual daily maintenance dose and plasma concentrations of theophylline should be maintened. No dosage adjustment is required for LUVOX Tablets. Warfania: When fluvoxamine malente (50 mg tid) was administered concentratinty with worfarin for two weeks, warfanin plasma concentrations increased by 98% and protinombin times were prolonged. Thus patients receiving and anticogulants and LUVOX Tablets should have their protinombin time monitored and their anticogulant dose adjusted accordingly. No dosage adjustment is required for LUVOX Tablets.

PRECAUTIONS

General Activation of Mania/Hypomania: During premarketing studies involving primarily depressed patients, hypomania or mania occurred in approximately 3% of patients treated with fluvoxamine. Activation of mania/hypomania has also been reported in a small proportion of patients with major affective disorder who were treated with fluvoxamine. Activation of mania/hypomania has also been reported in a small proportion of patients with major affective disorder who were treated with other marketed antidepressants. As with all antidepressants, LIVOX Tablets should be used cautiously in patients with a continuity of sectures. It should be discontinued in any patient who develops salvares. Surfacide: The possibility of saudies aftering its inherent in patients with degreesive symptoms, whether these occur in primary depression or in association with another primary desorder such as QCD. Close supervision of high risk patients should accompany initial drug therapy. Prescriptions for LIVOX Tablets should be written for the smallest quantity of tablets consistent with good patient management in order to reduce the risk of overdose. Use In Patients with Concomitant Illness: Closely monitored clinical experience with LIVOX Tablets have not been evaluated or used to be a patient with diseases or candidations that could direct hemodynamic responses or metablosis. LIVOX Tablets have not been evaluated or used be any appreciable extent in patients with a recent history of myocardial infarction or unstable heart disease. Patients with these diagnoses were systematically excluded from many clinical studies during the product's permarketing lesting. Evaluation of the electrocardiograms for patients with depression or CIV and participated in premarketing studies exceeded to differences between fluvoxorimal on the entergence of clinically important. EG changes. In patients with they 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 Physicians are advised to discuss the following suses with potents for whom they prescribe UVVX fables: Interference with Cognitive or More Performance: Since any psychodized dup may import judgement, historian, or moter skills, potents should be countioned about operating hazardous mochinery, including automobiles, until they are certain that LUVOX Tables therapy does not adversely affect their ability to engage in such activities. Nursing: Potents receiving LUVOX fables should be advised to notify their physicians if they are breast feeding an infram. Cee PRECAUTIONS - Nursing: Mothers.) Concentinant Medications: Patients should be advised to notify their physicians if they are toking, or plan to take, any prescription or overhercounter drugs, since there is a potential for clinically important interactions with LUVOX Tables. Alcohol: As with other psychotropic medications, patients should be advised to avoid alcohol while toking LUVOX Tables. Allorgic Reactions: Patients should be advised to notify their physicians if they are toking, or leaded allergic phenomenon during therapy with LUVOX Tables. Patients should be advised to notify their physicians if they are toking, or a related allergic phenomenon during therapy with LUVOX Tables. Alcohol: As with other psychotropic medications are called allergic phenomenon during therapy with LUVOX Tables.

Laboratory Tests

There are no specific laboratory tests recommended.

Drug Interactions

Drug Interactions
There have been rare postmarketing reports describing patients with weakness, hypereflexia, and incoordination following the use of a selective serotonin reuptake inhibitor (SSR) and sumotifyion. If concomitant heatment with sumatriptan and an SSR (e.g., fluosetine, fluovoamine, parasetine, sentraline) is clinically warranted, appropriate observation of the potient is advised, Potential Interactions with drugs that inhibit or are Metabolized by Cytachrome PSOS (posymens; Sessed on a finding of substantial interactions of throwoamine with certain drugs sand inimited in with of the IIII.44 isoenzyme, it appears that fluovoamine inhibits isoenzymes that are known to be involved in the metabolism of drugs such as wordarin, theophylline and proporation). A clinically significant fluovoamine inhibits isoenzymes that are known to be involved in the metabolism of drugs such as wordarin, theophylline and proporation). A clinically significant fluovoamine inhibits isoenzymes that are known to be involved in the metabolism of drugs such as wordarin, theophylline and proporation). A clinically significant fluovoamine inhibits isoenzymes that are known to be involved in the metabolism of drugs such as wordarin, theophylline and proporation. A clinically significant fluovoamine inhibits isoenzymes that are known to be involved in the metabolism of drugs such as wordarin, theophylline and proporation in the metabolism of drugs such as wordarine, the proporation of the metabolism of the metabolism of the such as a transfer of the metabolism of the such as a transfer of the metabolism of the such as a transfer of the such as a melationism data has a matrix metapolari, window, plasmia review analy or pramtivolyopianic elects of the little day state of melations are interested in the manual standy state conditions are enclosed. Please see complete prescribing information for recommendations regarding CNS drugs such as monocomine oxidose inhibitors, olprazolam, diazepam, lorazepam, lithium, tryptophan, dazapine, alcohol, tricyclic antidepressants, carbamazepine, methodone; and other drugs such as theophylline, progranolol and other betrablockes, worthin, digoxin, and dilitazen. Effects of Smokking on Fluvoxamine Metabolism: Smokers had a 25% increase in the methodolism of throwsamine compared for nonsmakers. Electroconvulsive Therapy (ECT): There are no clinical studies establishing the benefits or risks of combined use of ECT and fluvoxamine molecute.

studies establishing the benefits or risks of combined use of ECI and throxomine maleate.

Carcinogenesis, Mutagenesis, Impairment of Fertility

Carcinogenesis: There is no evidence of carcinogenesis: The late or in the steady of the carcinogenesis: The late or late of the carcinogenesis: The late or late or

daily dose on a mg/m* usus; nou no ense, on imming personance, security of the pregnancy Pregnancy Frequency Frequen

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

Nursing Mothers
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 tribe into account the potential for serious odverse effects from exposure to fluvoxamine in the nursing infant as well as the potential benefits of LUYOX** (fluvoxamine maleate) Tablets therapy to the mother.

The efficacy of fluvoxamine maleate for the treatment of Obsessive Compulsive Disorder was demonstrated in a 10-week multicenter placebo controlled study with 120 outpatients upges 817. The adverse event profile observed in that study was generally similar to that observed in odult studies with fluvoxamine (see MOVERSE REACTIONS).

Decreased appetite and weight loss have been observed 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.

Verticative Use
Approximately 230 potents 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 potents. Other reported clinical experience has not identified differences in response between the elderly and younger
patients. However, the clearance of fluvoramine is decreased by about 50% in elderly compared to younger patients. Gee Pharmacokinetics under CLINICAL
PHARMACOGOTY, and greater sensitivity of some older individuals also cannot be ruled out. Consequently, LUVOX tablets should be slowly intended during initiation

ADVERSE REACTIONS

Associated with Discontinuation of Treatment

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

Adverse events in OCD Pediatric Population
In pediatric prients (N=57) treated with LUVOX® Tablets, 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 (N=63) were: abnormal thinking, cough increase, dysmeortheq, ecclymosis, emotional lobility, epistoxis, hyperkinesia, infection, monic reaction, rost, sinustis, and verified decrease.

Events for which the incidence in fluvoxomine molecule was equal to or less than the incidence in placebo (N=63) and involved two or more of the pediatric

study potients were: abdominal pain, abnormal dreams, fever, headache, nausea, nervousness, pain, pharyngitis and rhinitis, Incidence in Controlled Trials - Commonly Observed Adverse Events in Controlled Clinical Trials: LUVOX Tablets have been studied in controlled trials of OCD (n=320) and depression (n=1350). In general, adverse event rates were similar in the two data sets. The most commonly observed adverse 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: somnolence, insomnia, nenrousness, tremon, nausea, dyspepsia, anorexia, vomitina, abnormal ejaculation, astheria, and sweating. In a pool of two studies involving only patients with OCD, the following additional events were identified using the above rule: dry mouth, decreased libida, urinary frequency, anargosmia, rhinitis and taste perversion. Adverse Events Occurring at an Incidence of 1%: Table 2 enumerates adverse events that occurred of a frequency of 1% or more, and were more frequent than in the placeba group, among patients treated with LUVOX tablets in two short term placeba 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 hable shows the percentage of patients in each group who had at least one occurrence of an event at some time during their treatment. Reported adverse 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 course of usual medical practice where patient characteristics and other factors may differ from those that prevailed in the clinical trials. Similarly, the cited frequencies cannot be compared with figures obtained from other clinical investigations involving different treatments, success, and investigators. The other figures, however, do provide the prescribing physician with some basis for estimating the relative contribution of drug and non-drug flotors to the side-effect incidence role in the population studied. Adverse Events in OCD Placebo Controlled 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 rate compared to event rates in OCD and depression studies were Placebo Controlled Studies: The events in CUD studies with a two-fold decrease in rate compared to event rates in CUD and depression studies when dysphagin cand subhypoin (marstly burned vision), Additionally, there was an approximate 25% decrease in mase. The events in COD and depression studies were: asthenia, abnormal ejoculation (mostly delayed ejoculation), anxiety, infection, thinitis, anargasmia (in males), depression, libido decreased, pharyngitis, agination, impotence, myodonas/heitch, thinist, weight loss, leg cramps, myagilar and unimy retentina. These events are listed in order of decreasing rates in the OCD trials.

Vital Sign Changes

Comparisons of fluvoramine maleate and placebo groups in separate pools of short-term OCD and depression trials on (1) median change from baseline on various wital signs variables and on (2) incidence of patients meeting criteria for potentially important changes from baseline on various vital signs variables.

Laboratory Changes

Comparisors of Howarmine maleate and placebo groups in separate pools of short-term OCD and depression trials on (1) median change from boseline on various serum chemistry, hematology, and urinalysis variables and on (2) incidence of patients meeting criteria for potentially important changes from boseline on various serum chemistry, hematology, and urinalysis variables revealed no important differences between fluvoxamine maleate and placebo.

Debenier on various securic varieties, in the control of the contr

POPULATIONS COMBINED (fluvoxomine [n=892] vs. plocebo (n=778] by potients—percentoge): BODY AS WHOLE: Headache (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); Ashenic (14 vs. 6); Thi Syndrome (3 vs. 2); Crills (2 vs. 1); CARDIOVASCULAR: Polytations (3 vs. 2); DISESTIVE SYSTEM: Nascola (9 vs. 14); Diarrhea (11 vs. 7); Constipation (10 vs. 5); Syspepsia (10 vs. 5); Anoexia (6 vs. 2); Venntling (5 vs. 2); Flatilence (4 vs. 3); Joint Disorder (3 vs. 14); Syspepsia (2 vs. 6); Netrovice (2 vs. 6); Incomic (2 vs. 10); Application (2 vs. 12); Particle (4 vs. 6); Termor (5 vs. 1); Anoisty (5 vs. 3); Viscolidation (3 vs. 1) Hypertonic (2 vs. 1); Applican (2 vs. 1); Decreased Libido (2 vs. 1); Degrees (1 vs. 6); Termor (5 vs. 1); Anoisty (5 vs. 3); Viscolidation (3 vs. 1) Hypertonic (2 vs. 1); Applican (2 vs. 1); Decreased Libido (2 vs. 1); Degrees (1 vs. 6); Termor (5 vs. 1); Anoisty (5 vs. 3); Viscolidation (3 vs. 1); Mary (4 vs. 14); Degree (2 vs. 1); Termor (2 vs. 1); Applican (2 vs. 1); Degree (2 vs. 6); Degree (2 vs. 6); Degree (2 vs. 6); Degree (2 vs. 6); Degree (2 vs. 7); Anoigsamic (2 vs. 0); Unionary Retention (1 vs. 0); URGGENITIAL: Almornal Epicolition (4 vs. 1); Whom (2 vs. 6); Degree (2 vs. 1); Anoigsamic (2 vs. 0); Unionary Retention (1 vs. 0); URGGENITIAL: Almornal Epicolition (4 vs. 1); Viscon (2 vs. 1); Anoigsamic (2 v

collegations. In the tabulations which follow, a standard LOSIAKT states uncurrancy terminancy; may been used to see the second properties of COSIAKT term for an event was so general as to be uninformative, it was replaced with a more informative term. The frequencies presented, therefore, represent the proportion of the 2737 patient exposures to multiple doses of fluvoxamine moderate who experienced an event of the type cited on at least one accosion while receiving fluvoxamine moderate. All reported events are included in the list below, with the following exceptions; 1) those events affected and the list below. The following exceptions; 1) those events affected and the list below. The following exceptions; 2) those events affected and the list below. represent the proportion of the 2737 potient exposures to multiple doses of fluxocomine molene who experienced on event of the type cited on televal need occasion which receiving fluxocomine molene. All reported events are included in the list below, with the following exceptions: 1) those events of leady listed in Table 2, which tabulates incidence rates of common adverse experiences in placebo-controlled OCD and depression clinical trials, are excluded; 2) those events for which a duty cause was considered remote (e.g., neoplasia, gastrionitestinal cardiname, hetpes simplex, hetpes zoster, application site exception, and unitariated pregnancy or a omithed; and 3) events which were reported in only one patient and judged to not be potentially evisions are not included. It is important to emphasize that, offlorogh the events reported did occur during treatment with fluxoxamine malente, a causal relationship individual of interval interval to the properties and examinented in order of detections graphenory using the following definitions: frequent adverse events are those occurring between 1/100 and 1/1000 proferits, and rure adverse events are those occurring between 1/100 and 1/1000 proferits, and rure adverse events are those socriting in between the area of the event are those socriting between the event are those occurring in the share 1/100 patients, and as a Whole: Frequent accidental injury, malaise; infrequent affects and the area of the event are those socriting between the event area of the event area those socriting between the event area of the event area of the event area frequent. Condential injury, malaise; infrequent affects of the event area nxusp., paryngsmus, pasmactive purmonary assesse, pneumona. Skin: Intequent: acre, alopeda, dry skin, ezzema, exfolarite dematilis, fruunalois; sebortinea, skin discolaration, untraaria. Special Senses: Infrequent: accommodation abnormal, conjunctivitis, deafines, diplopia, dry eyes, ser pain, eye pain, mydrasis, offits media, pousomia, photophobia, taste lass, visual field defect; Rare: corneal ulcer, retinal detachment. Unagenital Systems: Infrequent: aurita, breast pain, cystifis, delayed menstruation), dysuria, female locatalori, hematuria, menopause, menorrhagia', metorrhagia', noturia, polyviar, premenstrust ayndrome; ulmary incontinence, urinary tract infection, urinary urgency, urination impaired, voginal hemorrhage', vaginitis'; Rare: kidney calculus, hematospermia', oliguria.

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

Non-U.S Postmarketing Reports

Voluntary reports of adverse events in patients taking LUYOX Tablets that have been received since market introduction and are of unknown causal relationship to LUYOX Tablets use include: toxic epidermal necrolysis, Stevens-Johnson syndrome, Henoch-Schoenlein purpura, bullous eruption, pringism, agranulocytosis, eneuropathy, aplastic nomeria, complyafic reaction, hyponatremia, acute renal failure, hepatitis, and severe akinesia with fever when fluxoxamine was co-administered with antipsychotic medication.

SVL343

CAUTION: Federal law prohibits dispensing without prescription.

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

Pharmacia & Upiohn

Solvay Pharmaceuticals

EFFECTIVE FIRST-LINE SSRI THERAPY FOR OCD...



EMERGING FROM THE PROFOUND ANXIETY OF OCD

Low incidence of agitation

• 2% vs 1% for placebo

Low incidence of sexual dysfunction'

 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 recommended¹

fluvoxamine maleate

25 mg TABLETS 50 mg & 100 mg SCORED TABLETS

AVAILABLE IN 25-mg TABLETS

The first SSRI

indicated for OCD

in children and

adolescents

*Parameters occurring ≥ 1% with fluvoxamine maleate.

Please see brief summary of prescribing information on adjacent page.