LUVOX® (fluvoxomine 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

UNION Tables are indicated for the treatment of absessions 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, asternizole, or cisopride with LUVOX Tablets is contraindicated (see WARNINGS and PRECAUTIONS).

LUYOX Tablets are contraindicated in patients with a history of hypersensitivity to fluvoxamine malea

WARNINGS

UVOX Tobles are contraindated in patients with a history of hypersensitivity to fluvoxomine molecule.

WARNINGS
In patients receiving emother serotenin reuptake inhibitor drug in combination with monoamine oxidase inhibitors (MAOIs), there have been reports of serious, sometimes fortal, reactions. Therefore, it is recommended that LUYOX" Toblets not be used in combination with a MAOI, or within 14 days of discontinuing treatment with a MAOI. In addition, after stopping LUYOX" Toblets, at least 2 weeks should be allowed before starting a MAOI.

Terfenodine, astemizole and disapride are all metabolized by the cytechrome P450IIIA4 isoenzyme. Increased plasma concentrations of terfenodine, astemizole and disapride cause OT prolongation and have been associated with torsacke depoints-type ventricular todhycardia, sometimes fartal. 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 terfenodine, astemizole, are disapride.

Other Potentially Important Drug Interactions
(Also see PRECAUTIONS - Drug Interactions)
(Also see PRE

PRECAUTIONS
General
Activation of Mania/Hypomania: During premarketing studies involving primarily depressed patients, Inpanancia or mania occurred in approximately 1% of patients heated with fluorosanine. Activation of mania/hypomania has also been reported in a small proportion of patients with major affective disorder who were heated with other marketed mitidepessants. As with all antidepessants. (AUVX Tablets should be used contiously in patients with a bistory of mania. Setzwers: Uniting permarketing studies, setzwers were reported in 0.2% of fluorosanine-heated patients. LUVOX Tablets should be used contiously in patients with a listory of setzwers. It should be discontinued in any patient who develops setzwers. Setcide: The possibility of a saided entempt inherent in patients with developes setzwers setzwers. Setcide: The possibility of a saided entempt inherent in patients with discontinued in any patient who develops setzwers. Setcide: The possibility of a saided entempt of biblets consistent with good potent monoperment in order to reduce the intex of overdors. LUVOX Tablets should be written for the smallest quantities to be a Patients with Concentituant Illness: Classily monitored clinical experience with LUVOX Tablets in patients with diseases or conditions that could affect hemodynamic responses or metabolism. LUVOX Tablets have not been evaluated or used to any appreciation extent in patients with a cream history of movacradi infraction or unstable heart disease. Patients with these diagnoses systematically excluded from many clinical studies during the product's premarketing testing. Evolution or the electrocardograms for potentism with depression or OLO who protricipated in premarketing studies exheeled to differences between fluorosamine and placeds in the emergence of clinically importation or or clinically importation or clinically importation or or clinically importation and in emergence or clinically importation or developed in the difference of the emergence of clinically importation of t

Physicians are advised to discuss the following issues with patients for whom they prescribe LUYOX Tablets: Interference with Cognitive or Motor Performance: Since any psychoactive drug may impair judgement, thinking, or motor skills, patients should be coulineed about operating hazardous mechinery, including automobiles, until they are certain that LUYOX Tablets therapy does not adversely affect their ability to engage in such activities. Preparacy: Tolerats should be odivised to notify their physicions if they become pregnant or intend to become prognant or intend to prognant or prognant or over the prognant of the prognan

Laboratory Tests
There are no specific laboratory tests recommended.

Drug Interactions There have been rare

There has pecific loboratory tests recommended.

Drug Interactions

These have been are postmaketing reports describing patients with weakness, hyperaffecia, and incoordination following the use of a selective serotation in supploke inhibitor (SSR) and summittation. If conconstraint healment with sumutiquitor and an SSRI (e.g., fluoretine, fluoroxamine, paraetine, sertaline) is clinically warranted, appropriate between the first software and interactions of throwcomine with drugs that inhibit or are Metabolized by Cytechrome P450 Issaymees: Bosed on a hading of substantial interactions of throwcomine with earth and using such as warrant, heaphylline and proporated. A clinically significant fluoroxamine inhibits interaction is possible with drugs howing a narrow therapeutic ratio such as refreshed expensional. A clinically significant fluoroxamine interaction is possible with drugs howing a narrow therapeutic ratio such as refreshed, establization, the certain bearcative pieces and period of the service of consideration of the service of consideration in the constitution of the service of the service of constitution of the service of the se

daily dose on a mg/m² basis) had no effect on mating performance, autonian or generate, or progressor, and the progressor of the vaxamine 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 matinamines. However, in other reproduction studies in which pregnant tost were dosed through evaning there was (1) on increase in pup mortality at third (seen at 80 mg/kg and obove but not at 20 mg/kg), and (2) decreases in postmatia pup weights (seen at 160 but not at 80 mg/kg) and survival (seen at 80 mg/kg and obove but not at 20 mg/kg), (losses 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 of losst some of these results likely occurred secondarily to maternal toxicity, the role of a direct drug effect on the letuses or puts could not be roled out. There are no adequate and well-combined studies in pregnant women. Fluxoxamine maleate should be used during pregnancy only if the potential benefit justifies the potential isk to the letus.

Labor and Delivery

The effect of fluvoxamine on lobor and delivery in humans is unknown.

Nurshag Morthers

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 exposue to fluvoxomine in the nursing infant as well as the potential benefits of LUYOX* nine maleate) Tablets therapy to the mother.

Pediatric Use

The efficacy of fluvoxamine molecte 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 fluvoxamine (see ADVERSE REACTIONS).

Decreased appetite and weight last have been observed in association with the use of fluvoxamine as well as other SSRIs. Consequently, regular maniforing https://doi.org/10.1016/j.com/

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Genature Use
Approximately 230 portients ponicipating in controlled premothering studies with LUVOX Toblets were 65 years of oge or over. No overall chierences in safety were observed between these pottents and younger potients. Other reported clinical experience has not identified differences in response between the elderly and younger potients. They or younger potients they were the clientance of thursournine is decreased by about 50% in elderly compared to younger potients (see Pharmacokinetics under 1474). They was a potient of the poti

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 i

Adverse events in OCD Pediatric Population
In pediatric potents (N=57) heated with LUNOX* Toblets, the overall profile of odverse events is similar to that seen in colul 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, dynamionthea, actlymass, emotional lability, episitoxis, hyperkinesia, infection, manic reaction, rosh, sirusilis, and weight decrease.

Includes position from cyclindos, encount bodiny, spisous, presentable, instance, make each probability that proceeding the expension of the probability of the proba controlled into 6 OCO (n=20) and depression (n=1550). In general, obverse event rutes were similar in the two data sets. The most controlled and of the controlled and the controlled an

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Virtid Sign Changes

Comparisons of floroxomine molecule and placebo groups in separate pools of short-term OCD and depression trials on (1) median change from baseline on various virtil signs variables and or (2) incidence of patients meeting arterial for potentially important changes from baseline on various virtal signs variables.

Leberatory 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 serum chemistry, hematology, and urinalysis variables and on (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 maleate and placebo.

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

2: TREATMENT-EMERGENT ADVERSE EVENT INCIDENCE RATES BY BODY SYSTEM IN OCD AND DEPRESSION

no important differences between fluoxocamine mislente and placebo.

Table 2: TREATMENT-EMERGENT ADVERSE EVENT INCIDENCE RATES BY BODY SYSTEM IN OCD AND DEPRESSION POPULATIONS COMBINED (throwamine [n=872] vs. placebo [n=778] by potients—percentage): BODY AS WHOLE: Headache (22 vs. 20); Ashberia (14 vs. 6); Flu Syndrome (3 vs. 2); Callis (2 vs. 1). CARDIOVASCULAR: Poliptions (3 vs. 2). DIGESTIVE SYSTEM. Rouses (40 vs. 1); Distribe (11 vs. 1); Constipation (10 vs. 8): Dispessiol (10 vs. 5); Potience (6 vs. 2); Venting (5 vs. 2); Platente (4 vs. 3); Booth Biosode (3 vs. 1); Dispribagia (2 vs. 1). NERVOUS SYSTEM: Sormolence (22 vs. 8); Insomino (21 vs. 10); Dy Mouth (14 vs. 10); Nervounness (12 vs. 5); Diszniess (11 vs. 6); Plemor (5 vs. 1); Anvely (5 vs. 3); Viscoditation (3 vs. 1); Hypertonia (2 vs. 1); Agustion (2 vs. 1); Depression (2 vs. 6); Ols Srimulation (2 vs. 1); Ashbergonia (2 vs. 1); Depression (2 vs. 6); Ols Srimulation (2 vs. 1); Ashbergonia (3 vs. 2); Unpotence (2 vs. 1); Anongsmin (2 vs. 0); Unionary Retention (1 vs. 0);
Versus for which fluoxocamine melater incidence was equal to or less than placebo are not listed in the table above, but include the following: obdominal pain, abnormal dearns, apopethe increase, back pinc, thesi pain, includes "contactive," "Tool dearns, mylagia, pain, poresthesia, polyprographic, posturally phylosension, pruntis, scal, himitis, fluits and himitis. "Includes "contactive," "Tool dearns, mylagia, pain, poresthesia, polyprographic, posturally phylosension, pruntis, scal, himitis, fluits and himitis." Includes "contactive," "Tool dearns, apopethe increase, back power, thesi pain, includes "contactive," "Tool dearns and on advesses," more discusses, "Morty feeling warm, bto, or flushed. "Mostly" "Burrea vision." "Mostly "deleyed ejecution." "Includes "contactive," "Tool dearns and on advesses," on "a "caise." "Mostly related and provides and included. It is important to emplosize that, ofthough the events reported did occur during freatment with fluvoxomine molecute has not been established. Events are further dossified within body system categories and enumerated in order of design flequency using the following definitions: frequent offeresses events are defined as those occurring on one or more accessions in at least 1/100 professing flequency using the following definitions: frequent deverse events are defined as those occurring on one or more accessions in at least 1/100 professing flequency using the following definitions: frequent development offeresses of the sevents are those occurring on one or more accessions in at least 1/100 professing flequency as a Whole: Frequent codestine large-generated development of the profession of the professi

Based on the number of females, ²Based on the number of males.

Non-US Postmarketing Reports

Voluntury reports of adverse events in patients taking LUVOX Tablets that have been received since market introduction and are of unknown ausail relationship to LUVOX Tablets us entitled: notice toxic epidermal necrolysis, Stevens-Johnson syndrame, Henoch-Schoenlein purpura, bullous eruption, pringism, agranulacytoss, neuropathy, aplastic amenia, anaphylactic reaction, hyponatremia, acute read failure, hepatifis, and severe advinests with lever when fluvoxamine was co-administered with antiosychotic medication.

SVL343

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

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

Pharmacia&Upjohn

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 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 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



AVAILABLE IN 25-mg TABLETS



*Parameters occurring ≥ 1% with fluvoxamine maleate.

Please see brief summary of prescribing information on adjacent page.