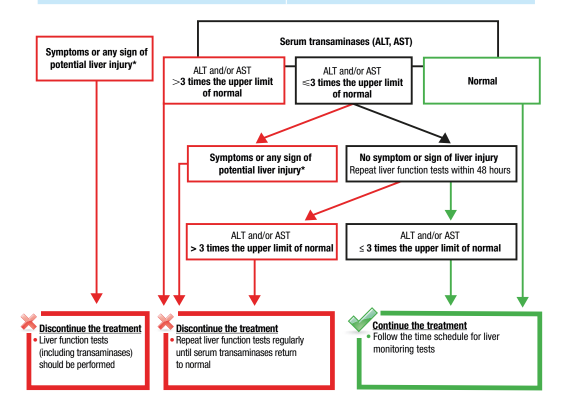
# Liver function monitoring scheme with Valdoxan

Patient name:

Date of initiation:

Valdoxan 25mg		In case of dose increase at restart the monitoring sche	50mg, me.
Before Initiation of 25mg	ALTU/L ASTU/L	Initiation of 50mg	ALTU/L ASTU/L
Week 3	ALTU/L ASTU/L	Week 3	ALTU/L ASTU/L
Week 6	ALTU/L ASTU/L	Week 6	ALTU/L ASTU/L
Week 12	ALTU/L ASTU/L	Week 12	ALTU/L ASTU/L
Week 24	ALTU/L ASTU/L	Week 24	ALTU/L ASTU/L
Please perform a test at any time if clinically justified.		Please perform a test at any time if clinically justified.	





... In the treatment of Major Depressive Episodes in Adults

# **INFORMATION FOR HEALTHCARE PROFESSIONALS**

# **Recommendations** regarding

- Liver function monitoring
- Interaction with potent CYP1A2 inhibitors

#### Valdoxan overview

• Valdoxan was registered in Europe in February 2009 and is available in Ireland since 01/06/2009 for the treatment of major depressive episodes in adults.

## Valdoxan and risk of hepatotoxicity

• Cases of liver injury, including hepatic failure (few cases were exceptionally reported with fatal outcome or liver transplantation in patients with hepatic risk factors), elevations of liver enzymes exceeding 10 times upper limit of normal, hepatitis and jaundice have been reported in patients treated with Valdoxan in the post-marketing setting. Most of them occurred during the first months of treatment. The pattern of liver damage is predominantly hepatocellular with serum transaminases which usually return to normal levels on cessation of Valdoxan.



PHYSICIAN'S GUIDE

\*Such as dark urine, light coloured stools, yellow skin/eyes, pain in the upper right belly, sustained new-onset and unexplained fatigue.

#### **Recommendations for liver function monitoring**

- Do not use Valdoxan in case of
  - Hepatic impairment (i.e. cirrhosis or active liver disease) or transaminases > 3ULN
- Before starting treatment
  - Caution for Valdoxan initiation in patients with hepatic injury risk factors

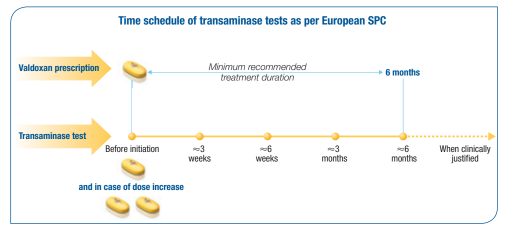
Valdoxan should be **prescribed after careful consideration of benefit and risk**:

- in patients with **hepatic injury risk factors** e.g. obesity/overweight/non-alcoholic fatty liver disease, diabetes, alcohol use disorder and /or substantial alcohol intake
- in patients receiving **concomitant** medicinal products associated with risk of hepatic injury.
- Checking patient's liver function tests

Baseline liver function tests should be **undertaken in all patients before starting treatment**:

- treatment should not be initiated in patients with baseline values of ALT and/or AST>3 ULN.
- caution should be in exercised patients with baseline values of ALT and/or AST > ULN and ≤ 3 ULN.

## • Prescribe transaminases tests (ALT/AST) for your patients



When increasing the dosage, liver function tests should again be performed at the same frequency as when initiating treatment.

Any patient who develops increased serum transaminases should have his/her liver function tests repeated within 48 hours.

Please see **Liver Monitoring Scheme with Valdoxan** at the back of this Guide to assist you with this.

• During treatment period

Valdoxan treatment **should be discontinued** immediately if:

- patient develops symptoms or signs of potential liver injury (such as **dark urine**, **light-coloured stools**, **yellow skin/eyes**, **pain in the upper right belly**, **sustained new-onset and unexplained fatigue**),
- the increase in serum transaminases exceeds 3X ULN.

Following discontinuation of Valdoxan therapy liver function tests should be repeated until serum transaminases return to normal.

#### Inform your patient about :

- the importance of liver function monitoring and,
- the vigilance about signs and symptoms of liver injury.

#### Reminder :

#### What to do in case of :

ALT and /or AST increase $\leq$ 3 ULN	Repeat the test within 48h	
ALT and/or AST increase > 3 ULN	Stop the treatment immediately , repeat the blood tests until normalization	
Signs and symptoms of liver injury *	Stop the treatment immediately, repeat the blood tests until normalization	

\* dark urine, light coloured stools, yellow skin/eyes, pain in the upper right belly, sustained new-onset and unexplained fatigue

## Interaction with potent CYP1A2 inhibitor

- Valdoxan is contraindicated with concomitant use of potent CYP1A2 inhibitors (e.g. fluvoxamine [Faverin®], ciprofloxacin [Ciproxin®]).
- Agomelatine is metabolised mainly by cytochrome P450 1A2 (CYP1A2) (90%) and by CYP2C9/19 (10%). Medicines that interact with these isoenzymes may decrease or increase the bioavailability of agomelatine. Fluvoxamine, a potent CYP1A2 and moderate CYP2C9 inhibitor, markedly inhibits the metabolism of agomelatine resulting in an increase in agomelatine exposure.
- In vivo, agomelatine does not induce CYP450 isoenzymes. Agomelatine inhibits neither CYP1A2 in vivo nor the other CYP450 in vitro. Therefore, Valdoxan is not expected to modify exposure to medicinal products metabolised by CYP450.

Please see the New Summary Product Characteristics and New Patient Information Leaflet at <u>www.medicines.ie</u>

Further copies of the Prescriber Guide or Patient Booklet may be obtained from the Medical Department at Servier Laboratories (Ireland) Ltd., 2nd Floor, 19 Lower Georges Street, Dun Laoghaire, Co. Dublin. Tel: 01-6638110 October 2021