

# MULTAQ<sup>®</sup> PRESCRIBER GUIDE

*This guide contains important safety information for the safe use of dronedarone (Multaq<sup>®</sup>)*

## Aim of this Guide:

To provide Multaq<sup>®</sup> (dronedarone) prescribers with a guide to:

1. Screen patients before treatment initiation
2. Monitor patients during treatment
3. Discontinue Multaq<sup>®</sup> when required
4. Counsel patients about its use

This is additional to the [Summary of Product Characteristics \(SmPC\)](#) and Patient Information Leaflet. Thus, it does not include the full prescribing information.

## Safe Use:

- Treatment with Multaq<sup>®</sup> should only be:
  - Initiated and monitored under specialist supervision
  - Prescribed after alternative treatment options have been considered
- Treatment with Multaq<sup>®</sup> can be initiated in an outpatient setting.

### Call for Reporting

Healthcare professionals should report any adverse events suspected to be associated with the use of Multaq<sup>®</sup> to Sanofi Malta Ltd., 3rd Floor, Avantech Building, St Julian's Road, San Gwann SGN 2805. Tel: 2149 3022, Fax: 2149 3024.

**Alternatively any suspected ADRS and medication error** can be reported to the Medicines Authority. Report Forms can be downloaded from [www.medicinesauthority.gov.mt/adrportal](http://www.medicinesauthority.gov.mt/adrportal) and posted to Medicines Authority, Post-Licensing Directorate, Sir Temi Zammit Building, Malta Life Sciences Park, San Gwann SGN 3000 or sent by email [postlicensing.medicinesauthority@gov.mt](mailto:postlicensing.medicinesauthority@gov.mt)

## BEFORE TREATMENT INITIATION

If **any** of the 'Yes' criteria (**Red Buttons**) apply, do not prescribe Multaq®. You should only prescribe Multaq® if **all** 'No' criteria (**Green Buttons**) apply. Contraindications should be confirmed by **ECG, serum creatinine and, liver and pulmonary tests**.

Multaq® is **indicated** for the maintenance of sinus rhythm after successful cardioversion in clinically stable adult patients with paroxysmal or persistent atrial fibrillation (AF)

Atrial Fibrillation

Yes

Permanent AF with an AF duration ≥6 months (or duration unknown) and attempts to restore sinus rhythm no longer considered by the physician

No

Heart Failure

Yes

History of, or current heart failure or left ventricular systolic dysfunction (LVSD)

Yes

Unstable hemodynamic conditions

Yes

Pre-renal azotaemia (functional impairment)

No

Drug - Drug Interactions

Yes

Potential torsades de pointes inducers (*phenothiazines, cisapride, bepridil, tricyclic antidepressants, terfenadine and certain oral macrolides*)

Yes

Potent cytochrome P 450 (CYP) 3A4 inhibitors (*ketoconazole, itraconazole, voriconazole, posaconazole, telithromycin, clarithromycin, nefazodone and ritonavir*)

Yes

Class I or Class III antiarrhythmics

Yes

Dabigatran

No

Liver & Lung

Yes

Severe hepatic impairment

Yes

Liver and lung toxicity related to the previous use of amiodarone

No

Kidney

Yes

Severe renal impairment (CrCl <30 ml/min)

No

Multaq® can be initiated

## MONITORING DURING TREATMENT

The following assessments are recommended during treatment with Multaq®. Criteria for discontinuation are also described. If **any** of the 'Yes' criteria (**Red Buttons**) arise during treatment, Multaq® should be discontinued.

### ECG:

Serially, at least every 6 months

Patient develops permanent AF

Yes

### SYMPTOMS OF:

- Heart failure
- LVSD (monitoring of left ventricular function)

Patient develops heart failure or LVSD

Yes

### USE WITH CAUTION (in association with):

- Digitalis
- Beta blockers, calcium antagonists with heart rate lowering properties, statins
- Drug modifying INR (warfarin)
- Sirolimus and tacrolimus

### NOT RECOMMENDED (in association with):

Grapefruit juice, potent CYP3A4 inducers including rifampicin, phenobarbital, carbamazepine, phenytoin, St John's Wort

### LIVER FUNCTION TESTS:

After 1 week → after 1 month → monthly for 6 months → at months 9 and 12 → periodically

ALT levels are confirmed to be ≥3 ULN

Yes

### PULMONARY FUNCTION TESTS

Pulmonary toxicity

Yes

**SERUM CREATININE\***: After 1 week → after a further 7 days if ↑ creatinine

Serum creatinine continues to ↑

Yes

\*Plasma creatinine levels may rise initially due to inhibition of renal tubular excretion of creatinine and are not necessarily indicative of a deterioration in renal function

## PATIENT COUNSELLING

Patients should be informed that during treatment with Multaq® **blood tests and ECGs** will be performed, and should be advised on the following:

**To consult a physician** if they develop: *palpitations, sensation of rapid or irregular heart beats*

**To consult a physician** if they develop: *weight gain, dependent oedema, increased dyspnoea*

Multaq® interacts with a number of medicines:

- **To inform any other doctor** that they are under treatment with Multaq®
- They **should not take** St. John's Wort
- They should **avoid** grapefruit juice

**To report immediately** if they develop: *new-onset abdominal pain, anorexia, nausea, vomiting, fever, malaise, fatigue, jaundice, dark urine or itching*

**To consult a physician** if they develop: *non-productive cough, breathlessness*

Reporting **suspected adverse drug reactions** is important for continued monitoring of the benefit/risk balance. Healthcare professionals are asked to report any suspected adverse reactions to Sanofi Malta Ltd via EMAIL: [PharmacovigilanceMalta@sanofi.com](mailto:PharmacovigilanceMalta@sanofi.com)