MULTAQ® PRESCRIBER GUIDE

This guide contains important safety information for the safe use of dronedarone (Multaq®)

Aim of this Guide:

To provide Multaq® (dronedarone) prescribers with a guide to:

- 1. Screen patients before treatment initiation
- 2. Monitor patients during treatment
- 3. Discontinue Multaq® when required
- Counsel patients about its use

This is additional to the <u>Summary of Product Characteristics (SmPC)</u> and Patient Information Leaflet. Thus, it does not include the full prescribing information.

Safe Use:

- Treatment with Multag® should only be:
 - o Initiated and monitored under specialist supervision
 - o Prescribed after alternative treatment options have been considered
- Treatment with Multaq® 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® 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 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



