

EU Declaration of Conformity

We, OROLIA Ltd., PB3 5PB Portsmouth – UK declare under our sole responsibility that the following product:

Product designation: **McMurdo Smartfind S5 AIS-SART
Kannad Marine Safelink AIS-SART**

Type or Model: **AIS Search And Rescue Locating Device**

Intended usage: these products meet SOLAS carriage requirements for an AIS Search And Rescue Locating Device, as specified in MED 4.55 Search and rescue locating devices (SRLD): AIS SART equipment. These products provide for on-scene Search And Rescue locating in maritime distress situations.

to which this declaration relates, has been tested and found to be in compliance with the essential requirements of Council Directive 2014/90/EU on marine equipment as indicated by Commission Implementing Regulation (EU) 2019-1397.

These products have been tested to verify compliance with the standards and requirements of the Directive as detailed in the Annex to this declaration.

This document reference N°: DOC20006A date 30/01/2020

These products carry the conformity marking:
(‘XX’ indicates year of manufacture)



**2443
XX**

Place and date of issue: Portsmouth, January 30th, 2020

Signed by or for the manufacturer:

Name: **Olivier WALSER**
Title: **EMEA PNT Beacons Quality Director**



Annex: Conformity Assessment

Standards and other normative documents

These products have been tested to verify compliance with all applicable requirements of the following international instruments, regulations and testing standards:

IEC 60945 (2002) incl. Corrigendum 1 (2008)
IEC 61097-14 (2010)
IMO Resolution A.694(17)
IMO Resolution MSC.246(83)
IMO Resolution MSC.302(87)
ITU-R M.1371-5 (2014)

MED conformity assessment modules

Conformity assessment modules B and D defined in the MED Directive have been applied to these products as detailed below:

EC TYPE EXAMINATION (MODULE B)

These products conform to BABT-MED000034 issued by TÜV SÜD BABT Octagon House, Concorde Way, Segensworth North, Fareham, Hampshire, PO15 5RL, United Kingdom -- Notified Body ID: 0168.

PRODUCTION QUALITY ASSURANCE (MODULE D)

These products are manufactured under DK-MED005707-H1 issued by TÜV SÜD Denmark -- Tuborg Boulevard 12, 3rd floor -- DK-2900 Hellerup -- Notified Body ID: 2443.

Technical Construction file held by:

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