EC DECLARATION OF CONFORMITY

ACR Electronics, Inc. hereby declares under its sole responsibility that the following product is in conformity with Directive 2014/90/EU of the European Parliament and of the Council of 23 July 2016 on Marine Equipment, and has been type examined as follows and has been assessed in accordance with MED/5.6 of EU implementing regulation 2017/306. In accordance with the relevant community harmonized legislation, the product will be marked with the MED Mark of Conformity as follows:

yy = Last two digits of the year in which the mark is affixed

ACR Electronics, Inc. hereby declares under its sole responsibility that the following product is in conformity with Directive 2011/65/EU of the European Parliament and of the Council of 8 June 2011 on the restriction of the use of certain hazardous substances in electrical and electronic equipment (RoHS2). In accordance with the relevant community harmonized legislation, the product will be marked with the CE conformity marking as follows:

Product:
406 MHz (COSPAS-SARSAT) Emergency Position-Indicating Radio Beacon (EPIRB)
MED Item A.1/5.6
Trade Name: GlobalFix™ V4
Model: RLB-41

Notified Body:
TÜV SÜD BABT, Notified Body No. 0168
Octagon House, Concorde Way, Fareham, Hampshire, PO15 5RL, UK
Type Examination (Module B) Certificate No.: BABT-MED000086 Issue:01
EC Quality System (Module D) Certificate No.: BABT-MED005722-H1 Issue 03

Regulations and Standards:
IEC 61097-2, Ed. 3.0, 2008
IMO Resolution A.810(19)
IMO Resolution A.694(17)
IMO Resolution A.662(16)
IMO Resolution MSC.56(66)
IMO Resolution MSC.120(74)
IMO MSC Circ.862
C/S T.001 (Nov. 2007)
C/S T.007 (Nov. 2007)

Manufacturer and holder of technical documents:
ACR Electronics Inc.
5757 Ravenswood Road
Fort Lauderdale, FL 33312 USA

Signed for and on behalf of ACR Electronics Inc.

Name: Dan Stankovic
Title: Director, Certification and Test
Date: March 16, 2017

This Declaration complies with ISO/IEC 17050-1:2004 and EU decision 768/2008/EC, Annex III
Notice to Distributors:

Please note that new MED requirements (MED MkII) call for the inclusion of the full Declaration of Conformity (see reverse) with all shipments of MED approved products. To remain in compliance, all distributors are now required to copy the Declaration of Conformity seen on the reverse of this page and include this copy with all onward shipments of applicable products. If this bulk master crate is not going to be broken down into smaller shipments, it is not necessary to make a copy of the Declaration of Conformity.